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=> file zcaplus
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FILE COVERS 1907 - 28 Sep 2007 VOL 147 ISS 15 FILE LAST UPDATED: 27 Sep 2007 (20070927/ED)

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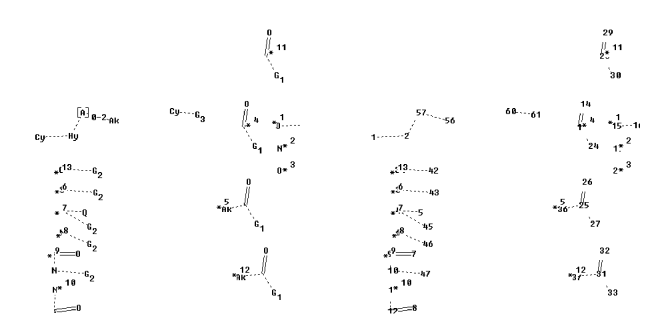
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=> d stat que L88 L12 775523 SEA FILE=REGISTRY ABB=ON PLU=ON N2C3/ES OR NOC3/ES L13 30896 SEA FILE=REGISTRY ABB=ON PLU=ON NSC3/ES L14 805906 SEA FILE=REGISTRY ABB=ON PLU=ON L12 OR L13 L15 464 SEA FILE=REGISTRY ABB=ON PLU=ON NPC3/ES L16 806370 SEA FILE=REGISTRY ABB=ON PLU=ON (L13 OR L14 OR L15) L19 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation: Uploading L19b.str



chain nodes :

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 20 24 25 26 27 28 29 30 31 32 33 34 36 37 42 43 45 46 47 48 56 57 60 61

ring/chain nodes :

16 17

chain bonds :

15-16 25-26 25-27 25-36 28-29 28-30 31-32 31-33 31-37 34-42 56-57 60-61 exact/norm bonds:

1-2 2-57 3-43 4-5 4-45 6-46 7-9 8-12 9-10 10-47 11-12 12-48 13-14 13-24

15-16 25-26 25-27 25-36 28-29 28-30 31-32 31-33 31-37 34-42 56-57 60-61

G1:[*1],[*2],[*3]

G2:[*4],[*5]

G3:[*6],[*7],[*8],[*9],[*10],[*11],[*12],[*13]

Connectivity:

2:2 M minimum RC ring/chain 20:1 E exact RC ring/chain

Match level :

1:Atom 2:Atom 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS 10:CLASS

11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS 20:CLASS

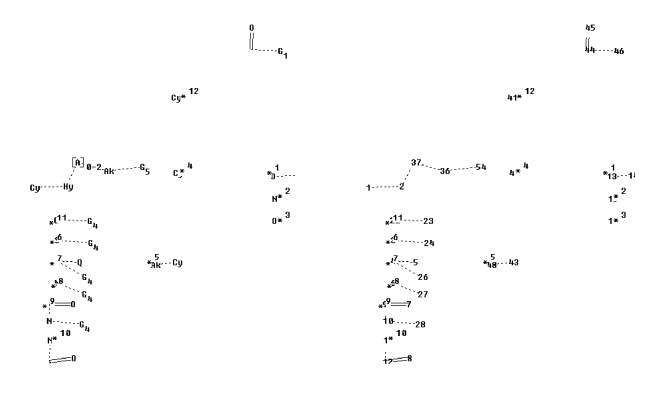
24:CLASS 25:CLASS

26:CLASS 27:CLASS 28:CLASS 29:CLASS 30:CLASS 31:CLASS 32:CLASS 33:CLASS 34:CLASS 36:CLASS 37:CLASS 42:CLASS 43:CLASS 45:CLASS 46:CLASS 47:CLASS 48:CLASS 56:CLASS 57:CLASS 60:Atom 61:CLASS Generic attributes : 60: Saturation : Unsaturated Type of Ring System : Monocyclic Element Count : Node 2: Limited N, N1-20,00-1 S,S0-1 P, P0-1C, C3

L21 71084 SEA FILE=REGISTRY SUB=L16 SSS FUL L19 L23 STR

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

Structure attributes must be viewed using STN Express query preparation: Uploading L23b.str



chain nodes : $1 \quad 2 \quad 3 \quad 4 \quad 5 \quad 6 \quad 7 \quad 8 \quad 9 \quad 10 \quad 11 \quad 12 \quad 13 \quad 18 \quad 22 \quad 23 \quad 24 \quad 26 \quad 27 \quad 28 \quad 29 \quad 36 \quad 37$ 41 42 43 44 45 46 48 54

```
ring/chain nodes :
14 15
chain bonds :
1-2 \quad 2-37 \quad 3-24 \quad 4-5 \quad 4-26 \quad 6-27 \quad 7-9 \quad 8-12 \quad 9-10 \quad 10-28 \quad 11-12 \quad 12-29 \quad 13-14 \quad 22-19 \quad 13-14 \quad 23-19 \quad 13-19 \quad 
36-37 36-54 43-48 44-45 44-46
exact/norm bonds :
1-2 \quad 2-37 \quad 3-24 \quad 4-5 \quad 4-26 \quad 6-27 \quad 7-9 \quad 8-12 \quad 9-10 \quad 10-28 \quad 11-12 \quad 12-29 \quad 13-14 \quad 22-19 \quad 13-14 \quad 23-19 \quad 13-14 \quad 23-19 \quad 13-19 \quad 
36-37 36-54 43-48 44-45 44-46
G1:[*1],[*2],[*3]
G4:[*4],[*5]
G5:[*6],[*7],[*8],[*9],[*10],[*11],[*12]
Connectivity:
2:2 M minimum RC ring/chain 18:1 E exact RC ring/chain
Match level:
1:Atom 2:Atom 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS
10:CLASS
11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 18:CLASS 22:CLASS 23:CLASS
24:CLASS 26:CLASS
27:CLASS 28:CLASS 29:CLASS 36:CLASS 37:CLASS 41:Atom 42:Atom 43:Atom
44:CLASS 45:CLASS
46:CLASS 48:CLASS 54:CLASS
Generic attributes :
41:
Saturation
                                                                                                                                                           : Unsaturated
Type of Ring System
                                                                                                                                           : Monocyclic
 42:
                                                                                                                                                           : Unsaturated
Saturation
Type of Ring System
                                                                                                                                                     : Monocyclic
43:
Saturation
                                                                                                                                                            : Unsaturated
                                                                                                                                            : Monocyclic
Type of Ring System
Element Count :
Node 2: Limited
                         N, N1-2
                           0.00-1
                           S.S0-1
                         P,P0-1
                           C,C3
```

```
L25 31522 SEA FILE=REGISTRY SUB=L21 SSS FUL L23 L29 STR
```

Structure diagram not available for display

Structure attributes must be viewed using STN Express query preparation: Uploading L29b.str

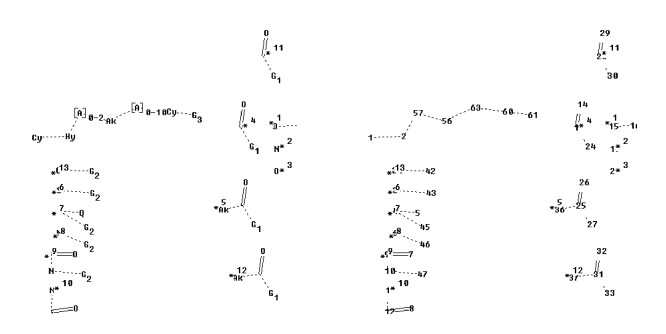
```
42
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            41---- ьз
                                                                                                                                                                                           Cj* <sup>12</sup>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             3{* <sup>12</sup>
                                                                                                                                                                                          C!* 4
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             3'* <sup>4</sup>
                                                                                                                                   *13....-G5
                                                                                                                                                                                                                                                                                                                                                                                                                                     *36-----51
                                                                           cy-----Hy------G
                                                                                                                                                                                                                                                                                                                                                                           1-----83
                                                                                                                                                                                                                                                                          N* 2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              1.* <sup>2</sup>
                                  *_0^{14}-Ak<sup>--G</sup>5 *_{c}^{11}-....<sub>G4</sub>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             1* 3
                                                                                                                                                                                                                                                                                                                                    *52°-63--64
                                                                                                                                                                                                                                                                            0*3
                                                                                                                                                                                                                                                                                                                                                                                           *211----23
                                 *5---Ak** G5
                                                                                                                                                                                                                                                                                                                                   *58--65~-66
                                                                                          **6.....24
                                     .∗′-16<sup>k--6</sup>5
                                                                                                                                                                                                                                                                                                                                     167--68
5*--55
                                                                                                                                                                                                                                                                                                                                                                                             *1<sup>7</sup><\_5
                                                                                           * <sup>7</sup>∴0
                                                                                                                                                                         *<sub>AR</sub>- --Cy
                                                                                                                                                                                                                                                                                                                                                                                                                                                                           *45--40
                                                                                         *\s\___G4
                                                                                                                                                                                                                                                                                                                                                                                            *<sup>58</sup> - 27
                                  *N---AK--G5
                                                                                                                                                                                                                                                                                                                                    *56--69--70
                                 * <u>18</u>=0
                                                                                                                                                                                                                                                                                                                                    *<sup>5</sup>′-≕61
                                                                                     <u>.9</u>___n
                                                                                                                                                                                                                                                                                                                                                                                        *§<sup>9</sup>—7
                                                                                                                                                                                                                                                                                                                                        57--71--72
                                                                                                                                                                                                                                                                                                                                                                                         10----28
                                      N-Ak G5
                                                                                                                                                                                                                                                                                                                                                                                         1* <sup>10</sup>
                                  *I.19
                                                                                                                                                                                                                                                                                                                                    <sub>*1</sub>19
                                                                                                                                                                                                                                                                                                                                                                                           12===8
                                                                                                                                                                                                                                                                                                                                        66=62
                                          ==0
                                             Ak--Gc
                                                                                                                                                                                                                                                                                                                                               79--74
chain nodes :
1 \quad 2 \quad 3 \quad 4 \quad 5 \quad 6 \quad 7 \quad 8 \quad 9 \quad 10 \quad 11 \quad 12 \quad 13 \quad 18 \quad 22 \quad 23 \quad 24 \quad 26 \quad 27 \quad 28 \quad 29 \quad 36 \quad 38
39 40 41 42 43 45 51 52 53 54 55 56 57 58 59 60 61 62 63 64 65
66 67 68 69
70 71 72 73 74
                                                                                                                                      8.3
ring/chain nodes :
14 15
chain bonds :
1-2 \quad 2-83 \quad 3-24 \quad 4-5 \quad 4-26 \quad 6-27 \quad 7-9 \quad 8-12 \quad 9-10 \quad 10-28 \quad 11-12 \quad 12-29 \quad 13-14 \quad 22-19 \quad 13-14 \quad 23-19 \quad 13-19 \quad 
23
36-51 40-45 41-42 41-43 52-63 53-65 54-55 54-67 56-69 57-59 57-71 58-60
59-61 60-62
60-73 63-64 65-66 67-68 69-70 71-72 73-74
exact/norm bonds :
1-2 \quad 2-83 \quad 3-24 \quad 4-5 \quad 4-26 \quad 6-27 \quad 7-9 \quad 8-12 \quad 9-10 \quad 10-28 \quad 11-12 \quad 12-29 \quad 13-14 \quad 22-19 \quad 13-14 \quad 23-19 \quad 13-14 \quad 23-19 \quad 13-14 \quad 23-19 \quad 13-19 \quad 
23
36-51 40-45 41-42 41-43 52-63 53-65 54-55 54-67 56-69 57-59 57-71 58-60
59-61 60-62
60-73 63-64 65-66 67-68 69-70 71-72 73-74
G1:[*1],[*2],[*3]
G4:[*4],[*5]
G5:[*6],[*7],[*8],[*9],[*10],[*11],[*12]
G6:[*13],[*14],[*15],[*16],[*17],[*18],[*19]
```

```
Connectivity:
2:2 M minimum RC ring/chain 18:1 E exact RC ring/chain 36:2 E exact RC ring/chain
45:2 E exact RC ring/chain 63:2 E exact RC ring/chain 65:2 E exact RC ring/chain
67:2 E exact
RC ring/chain 69:2 E exact RC ring/chain 71:2 E exact RC ring/chain 73:2 E exact
RC ring/chain
Match level:
1:Atom 2:Atom 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS
10:CLASS
11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 18:CLASS 22:CLASS 23:CLASS
24:CLASS 26:CLASS
27:CLASS 28:CLASS 29:CLASS 36:CLASS 38:Atom 39:Atom 40:Atom 41:CLASS
42:CLASS 43:CLASS
45:CLASS 51:CLASS 52:CLASS 53:CLASS 54:CLASS 55:CLASS 56:CLASS 57:CLASS
58:CLASS 59:CLASS
60:CLASS 61:CLASS 62:CLASS 63:CLASS 64:CLASS 65:CLASS 66:CLASS 67:CLASS
68:CLASS 69:CLASS
70:CLASS 71:CLASS 72:CLASS 73:CLASS 74:CLASS 83:CLASS
Generic attributes :
38:
Saturation
                     : Unsaturated
Type of Ring System
                   : Monocyclic
39:
Saturation
                      : Unsaturated
Type of Ring System
                     : Monocyclic
40:
Saturation
                     : Unsaturated
Type of Ring System
                   : Monocyclic
Element Count:
Node 2: Limited
   N, N1-2
   0.00-1
   S,S0-1
   P, P0-1
   C,C3
```

Structure attributes must be viewed using STN Express query preparation: Uploading L46b.str

L43 16848 SEA FILE=REGISTRY SUB=L25 SSS FUL L29 L46 STR

 $^{^{\}star}$ STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *



chain nodes :

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 20 24 25 26 27 28 29 30 31 32 33 34 36 37 42 43 45 46 47 48 56 57 60 61 63 ring/chain nodes:

16 17

chain bonds :

exact/norm bonds :

G1:[*1],[*2],[*3]

G2:[*4],[*5]

G3:[*6],[*7],[*8],[*9],[*10],[*11],[*12],[*13]

Connectivity:

2:2 M minimum RC ring/chain 20:1 E exact RC ring/chain

Match level:

1:Atom 2:Atom 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS 10:CLASS

11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS 20:CLASS

24:CLASS 25:CLASS

26:CLASS 27:CLASS 28:CLASS 29:CLASS 30:CLASS 31:CLASS 32:CLASS 33:CLASS

```
34:CLASS
36:CLASS 37:CLASS 42:CLASS 43:CLASS 45:CLASS 46:CLASS 47:CLASS 48:CLASS 56:CLASS 57:CLASS 60:Atom 61:CLASS 63:CLASS Generic attributes:
60:
Saturation : Unsaturated Type of Ring System : Monocyclic

Element Count:
Node 2: Limited
N,N1-2
O,O0-1
S,S0-1
P,P0-1
C,C3
```

```
L48
         8395 SEA FILE=REGISTRY SUB=L43 SSS FUL L46
L49
         3169 SEA FILE=REGISTRY ABB=ON PLU=ON L48 AND NRS<4
         1312 SEA FILE=REGISTRY ABB=ON PLU=ON L49 AND 16.165.12/RID
L51
          785 SEA FILE=REGISTRY ABB=ON PLU=ON 16.167.5/RID AND L49
L57
          2091 SEA FILE=REGISTRY ABB=ON PLU=ON L51 OR L57
L58
L63
          5 SEA FILE=REGISTRY ABB=ON PLU=ON 16.171.9/RID AND L49
         2096 SEA FILE=REGISTRY ABB=ON PLU=ON L58 OR L63
L64
          383 SEA FILE=ZCAPLUS ABB=ON PLU=ON L64
          108 SEA FILE=ZCAPLUS ABB=ON PLU=ON L65 AND J/DT
L66
          275 SEA FILE=ZCAPLUS ABB=ON PLU=ON L65 AND P/DT
L67
           26 SEA FILE=ZCAPLUS ABB=ON PLU=ON L66 AND PY<2003
78 SEA FILE=ZCAPLUS ABB=ON PLU=ON L67 AND PD<20020524
L68
L69
L72
          104 SEA FILE=ZCAPLUS ABB=ON PLU=ON L68 OR L69
         1945 SEA FILE=ZCAPLUS ABB=ON PLU=ON MAEKAWA T?/AU
L76
L77
          497 SEA FILE=ZCAPLUS ABB=ON PLU=ON HARA R?/AU
          263 SEA FILE=ZCAPLUS ABB=ON PLU=ON ODAKA H?/AU
L78
         7435 SEA FILE=ZCAPLUS ABB=ON PLU=ON KIMURA H?/AU
L79
           14 SEA FILE=ZCAPLUS ABB=ON PLU=ON MIZUFUNE H?/AU
L80
           169 SEA FILE=ZCAPLUS ABB=ON PLU=ON FUKATSU K?/AU
L81
             2 SEA FILE=ZCAPLUS ABB=ON PLU=ON L72 AND (L76 OR L77 OR L78 OR
L82
               L79 OR L80 OR L81)
L83
            11 SEA FILE=ZCAPLUS ABB=ON PLU=ON L76 AND (L77 OR L78 OR L79 OR
               L80 OR L81)
             1 SEA FILE=ZCAPLUS ABB=ON PLU=ON L77 AND (L78 OR L79 OR L80 OR
L84
               L81)
L85
            15 SEA FILE=ZCAPLUS ABB=ON PLU=ON L78 AND (L79 OR L80 OR L81)
L86
            1 SEA FILE=ZCAPLUS ABB=ON PLU=ON L79 AND (L80 OR L81)
L87
            1 SEA FILE=ZCAPLUS ABB=ON PLU=ON L80 AND L81
L88
            20 SEA FILE=ZCAPLUS ABB=ON PLU=ON (L82 OR L83 OR L84 OR L85 OR
               L86 OR L87)
```

=> d ibib abs hitind L88 1-20

L88 ANSWER 1 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2006:392887 ZCAPLUS <u>Full-text</u> DOCUMENT NUMBER: 144:420976

TITLE: Steady state operation research in JT-60U with

AUTHOR(S):

extended pulse length Fujita, T.; Akasaka, H.; Akino, N.; Ando, T.; Anno, K.; Arai, T.; Asakura, N.; Ashikawa, N.; Azechi, H.; Azumi, M.; Bakhtiari, M.; Bruskin, L.; Chankin, A.; Cheng, C. Z.; Chiba, S.; Cho, T.; Chu, M.-S.; Ding, B. J.; Ebisawa, N.; Fujii, T.; Fujita, T.; Fukuda, T.; Fukuyama, A.; Funaba, H.; Furukawa, H.; Furukawa, M.; Gao, X.; Gohil, P.; Gorelenkov, N. N.; Gotoh, Y.; Grisham, L.; Haga, S.; Hamamatsu, K.; Hamano, T.; Hanada, K.; Hasegawa, K.; Hashizume, H.; Hatae, T.; Hatayama, A.; Hayashi, N.; Hayashi, T.; Higashijima, S.; Hino, T.; Hiranai, S.; Hirano, Y.; Hiratsuka, H.; Hirohata, Y.; Hobirk, J.; Honda, A.; Honda, Masao; Honda, Mitsuru; Horiike, H.; Hoshino, K.; Hosogane, N.; Hosoyama, H.; Ichige, H.; Ichimura, M.; Ida, K.; Ide, S.; Idehara, T.; Idei, H.; Idomura, Y.; Igarashi, K.; Iio, S.; Ikeda, Y.; Imai, T.; Inagaki, S.; Inoue, A.; Inoue, D.; Isayama, A.; Ishida, S.; Ishii, K.; Ishii, Y.; Ishikawa, M.; Ishimoto, Y.; Itami, K.; Itoh, Sanae; Itoh, Satoshi; Iwasaki, K.; Kajiwara, K.; Kajiyama, S.; Kakimoto, S.; Kamada, Y.; Kaminaga, A.; Kamiya, K.; Kashiwa, K.; Katayama, K.; Kato, T.; Kawai, M.; Kawamata, Y.; Kawano, Y.; Kawasaki, T.; Kawashima, H.; Kazawa, M.; Kikuchi, H.; Kikuchi, K.; Kikuchi, M.; Kimura, A.; Kimura, M.; Kishimoto, Y.; Kitamura, S.; Kitsunezaki, A.; Kiyono, K.; Kizu, K.; Kobatake, N.; Kobayashi, S.; Kobayashi, Y.; Kodama, K.; Koide, Y.; Kokubo, S.; Kokusen, S.; Komata, M.; Komori, A.; Kondoh, T.; Konishi, S.; Konoshima, S.; Konovalov, S.; Koyama, A.; Koyanagitsu, M.; Kubo, H.; Kubo, T.; Kudoh, Y.; Kurihara, K.; Kurita, G.; Kuriyama, M.; Kusama, Y.; Kusanagi, N.; Lao, L. L.; Lee, S.; Li, J.; Litaudon, X.; Loarte, A.; Lonnroth, J.; Luce, T.; Maekawa, T.; Masaki, K.; Matsuda, T.; Matsukawa, M.; Matsumoto, T.; Matsunaga, G.; Matsuoka, M.; Matsuzawa, Y.; Meguro, K.; Mikhailivskii, A.; Mima, K.; Mironov, M. I.; Mitarai, O.; Miura, Yukitoshi; Miura, Yushi; Miya, N.; Miyamoto, S.; Miyato, N.; Miyo, Y.; Mogaki, K.; Morimoto, Y.; Morioka, A.; Moriyama, S.; Murakami, M.; Nagami, M.; Nagasaka, Y.; Nagasaki, K.; Nagase, Y.; Nagaya, S.; Nagayama, Y.; Naito, O.; Nakajima, N.; Nakamura, K.; Nakamura, Y.; Nakano, T.; Nakashima, Y.; Nakatsuka, M.; Narushima, Y.; Nazikian, R.; Neudatchin, S. V.; Ninomiya, H.; Nishikawa, M.; Nishimura, K.; Nishino, N.; Nishitani, T.; Nishiyama, T.; Noda, N.; Noto, K.; Oasa, K.; Obuchi, T.; Ogawa, H.; Ogawa, I.; Ogawa, Y.; Ohga, T.; Ohno, N.; Ohshima, K.; Oikawa, A.; Oikawa, T.; Okabayashi, M.; Okamoto, N.; Okano, F.; Okano, J.; Okano, K.; Okuno, K.; Omori, S.; Omori, Y.; Onishi, A.; Ono, Y.; Oohara, H.; Oshima, T.; Oya, Y.; Oyama, N.; Ozeki, T.; Parail, V.; Peterson, B. J.; Porter, G. D.; Rewoldt, G.; Sagara, A.; Saibene, G.; Saito, T.; Sakamoto, M.; Sakamoto, Y.; Sakasai, A.; Sakata, S.; Sakuma, T.; Sakurai, S.; Sasajima, T.; Sasao, M.; Sato, F.; Sato, M.; Sawada, K.; Sawahata, M.; Seimiya, M.; Seki, M.; Sharpe, J. P.; Shibahara, T.; Shimada, K.; Shimada, R.; Shimizu, A.; Shimizu, K.; Shimizu, M.; Shimono, M.; Shinohara, K.; Shinozaki, S.; Shirai, H.; Shiraiwa, S.; Shitomi,

M.; Sudo, S.; Sueoka, M.; Sugawara, A.; Sugie, T.; Sugiyama, K.; Sunaoshi, H.; Suzuki, Masaei; Suzuki, Mitsuhiro; Suzuki, S.; Suzuki, T.; Suzuki, Yoshio; Suzuki, Yutaka; Takahashi, K.; Takahashi, M.; Takamura, S.; Takano, S.; Takase, Y.; Takechi, M.; Takei, N.; Takeishi, T.; Takenaga, H.; Takizuka, T.; Tamai, H.; Tamura, N.; Tanabe, T.; Tanai, Y.; Tanaka, J.; Tanaka, Satoru; Tanaka, Shigetoshi; Terakado, H.; Terakado, M.; Terakado, T.; Toi, K.; Tokuda, S.; Totsuka, T.; Toudo, Y.; Tsuchiya, K.; Tsugita, T.; Tsukahara, Y.; Tsuzuki, K.; Tuda, T.; Uda, T.; Ueda, Y.; Uehara, K.; Uehara, T.; Ueno, Y.; Uesugi, Y.; Umeda, N.; Urano, H.; Urata, K.; Ushigome, M.; Ushigusa, K.; Usui, K.; Wade, M.; Wakatani, M.; Wang, S.; Watari, T.; Yagi, M.; Yagi, Y.; Yagisawa, H.; Yaqyu, J.; Yamada, H.; Yamamoto, T.; Yamamoto, Y.; Yamashita, Y.; Yamazaki, H.; Yamazaki, K.; Yatsu, K.; Yokokura, K.; Yonekawa, I.; Yoshida, Hajime; Yoshida, Hidetoshi; Yoshida, Hidetsugu; Yoshida, M.; Yoshida, N.; Yoshikawa, A.; Zushi, H.

CORPORATE SOURCE:

PUBLISHER:

Naka Fusion Research Establishment, Japan Atomic Energy Research Institute, Naka, Ibaraki, 311-0193,

Japan

SOURCE: Nuclear Fusion (2006), 46(3), S3-S12

CODEN: NUFUAU; ISSN: 0029-5515 Institute of Physics Publishing

DOCUMENT TYPE: Journal LANGUAGE: English

AB Recent exptl. results for steady state operation research in JT-60U are presented with emphasis on extension of sustained duration of high performance. The duration of heating has been extended from 10 to 30 s, and plasma properties and dynamics have been investigated in a long time scale exceeding the current diffusion time and close to the wall saturation time on ELMy H-mode, high βp H-mode and reversed shear H-mode regimes. The duration of sustainment of high beta and/or a large fraction of bootstrap current has been extended. The particle control with the saturated wall has been studied. Development of real-time control of q profile and effects of toroidal rotation on ELMs and the QH-mode are also discussed.

CC 71-2 (Nuclear Technology)

REFERENCE COUNT: 42 THERE ARE 42 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 2 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2005:1243660 ZCAPLUS Full-text

DOCUMENT NUMBER: 143:467580

TITLE: Overview of JT-60U progress towards steady-state

advanced tokamak

AUTHOR(S): Ide, S.; Akasaka, H.; Akino, N.; Ando, T.; Anno, K.;

Arai, T.; Asakura, N.; Ashikawa, N.; Azechi,

H.; Azumi, M.; Bakhtiari, M.; Bruskin, L.; Chankin, A.; Cheng, C. Z.; Chiba, S.; Cho, T.; Chu, M.-S.; Ding, B. J.; Ebisawa, N.; Fujii, T.; Fujita, T.; Fukuda, T.; Fukuyama, A.; Funaba, H.; Furukawa, H.; Furukawa, M.; Gao, X.; Gohil, P.; Gorelenkov, N. N.; Gotoh, Y.; Grisham, L.; Haga, S.; Hamamatsu, K.; Hamano, T.; Hanada, K.; Hasegawa, K.; Hashizume, H.; Hatae, T.; Hatayama, A.; Hayashi, N.; Hayashi, T.; Higashijima, S.; Hino, T.; Hiranai, S.; Hirano, Y.; Hiratsuka, H.; Hirohata, Y.; Hobirk, J.; Honda, A.; Honda, Masao; Honda, Mitsuru; Horiike, H.; Hoshino,

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S.; Watari, T.; Yagi, M.; Yagi, Y.; Yagisawa, H.; Yagyu, J.; Yamada, H.; Yamamoto, T.; Yamamoto, Y.; Yamashita, Y.; Yamazaki, H.; Yamazaki, K.; Yatsu, K.; Yokokura, K.; Yonekawa, I.; Yoshida, Hajime; Yoshida, Hidetoshi; Yoshida, Hidetsugu; Yoshida, M.; Yoshida, N.; Yoshikawa, A.; Zushi, H.

CORPORATE SOURCE: Naka Fusion Research Establishment, Japan Atomic

Energy Research Institute, Naka, Ibaraki, 311-0193,

Japan

SOURCE: Nuclear Fusion (2005), 45(10), S48-S62

CODEN: NUFUAU; ISSN: 0029-5515

PUBLISHER: Institute of Physics Publishing

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

A review. Recent exptl. results from steady-state advanced tokamak (AT) AΒ research on JT-60U are presented with emphasis on time scales longer in comparison with the characteristic time scales in plasmas. To achieve this, modification of the controls for the operation, heating and diagnostics systems have been carried out. As a result, .apprx.60 s current flat top and a .apprx.30 s H-mode are obtained. The long pulse modification has opened a door into a new domain for JT-60U. High normalized beta (β N) of 2.3 is maintained for 22.3 s and 2.5 for 16.5 s in a high- βp H-mode plasma. A standard ELMy H-mode plasma has also been extended and changes in the wall recycling on the longer time scale have been unveiled. The development and investigation of plasmas relevant to AT operation have been continued in long discharges as well as in discharges where higher NB power is available (≤ 10 s). Higher βN (.apprx.3) is maintained for 6.2 s in a high- βp H-mode plasma. High bootstrap current fraction (fBS) of .apprx.75% is sustained for 7.4 s in a reversed shear plasma. Neo-classical tearing mode (NTM) suppression by localized ECCD is found to be more effective with ECRF injection preceding the mode saturation The mode is suppressed with less power compared to the injection after the mode sats. The domain of the NTM suppression expts. is extended to the high- βN regime, and the effectiveness of m/n = 3/2 mode suppression by ECCD is demonstrated at βN .apprx. 2.5-3. Genuine tokamak plasma start up without a central solenoid is demonstrated. In a current hole region, it is shown that no scheme drives current in any direction. Detailed measurement of energetic ions in both space and energy showed dynamic change in the energetic ion profile due to collective instabilities. The impact of toroidal plasma rotation on ELM behavior is clarified in the grassy ELM and QH domains. Retention of hydrogen isotopes in the divertor tiles is analyzed.

CC 71-0 (Nuclear Technology)

REFERENCE COUNT: 40 THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 3 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2004:519930 ZCAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER: 141:64840

TITLE: A novel oxyiminoalkanoic acid derivative, TAK-559, activates human peroxisome proliferator-activated

receptor subtypes

AUTHOR(S): Sakamoto, Junichi; Kimura, Biroyuki;

Moriyama, Shinji; Imoto, Hiroshi; Momose, Yu;

Odaka, Biroyuki; Sawada, Hidekazu

CORPORATE SOURCE: Pharmaceutical Discovery Center, Pharmaceutical

Research Division, Takeda Chemical Industries, Ltd.,

Osaka, Japan

SOURCE: European Journal of Pharmacology (2004), 495(1), 17-26

CODEN: EJPHAZ; ISSN: 0014-2999

PUBLISHER: Elsevier

DOCUMENT TYPE: Journal LANGUAGE: English

A novel oxymminoalkanoic acid derivative, TAK-559, (E)-4-[4-[(5-methy)-2phenyl-1, 3-oxazol-4-yl)methoxy]benzyloxyimino]-4-phenylbutyric acid, was synthesized as a candidate of a new type of insulin-sensitizing agent. We report here activation of human peroxisome proliferator-activated receptor (hPPAR) subtypes by TAK-559. In a transient transactivation assay, TAK-559 was a potent hPPAR γ 1 and hPPAR α agonist with EC50 values of 31 and 67 nM, resp. Furthermore, TAK-559 was a partial agonist for hPPARyl with about 68% of maximal activation obtained with rosiglitazone (5-(4-(2-(methyl(2pyridinyl)amino)ethoxy) benzyl)-1,3-thiazolidine-2,4-dione), a thiazolidinedione derivative, which is known as a PPARy agonist. PPAR δ was significantly activated at a high concentration (10 μM) of TAK-559. Competition-binding assays using radiolabeled ligand indicated that the transactivation of all hPPAR subtypes by TAK-559 was due to direct binding of TAK-559 to each subtype. We also demonstrated that TAK-559 acts to recruit the coactivator SRC-1 to each of hPPARy1 and hPPAR α , and to dissociate the corepressor NCoR from each of hPPARyl and hPPARlpha. Taken together, we conclude that TAK-559 is a dual agonist for hPPAR γ 1 and hPPAR α with nearly equal EC50 values, a partial agonist for hPPARy1, and has a rather slight agonist activity for hPPAR δ .

CC 1-10 (Pharmacology)

REFERENCE COUNT: 41 THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 4 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2004:252494 ZCAPLUS Full-text

DOCUMENT NUMBER: 140:287404

TITLE: Preparation of five-membered heterocyclic compounds

for treatment of obesity, diabetes, hyperlipidemia,

etc.

INVENTOR(S): Momose, Yu; Takakura, Nobuyuki; Maekawa,

Tsuyoshi; Odaka, Hiroyuki; Kimura,

Hiroyuki

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Japan

SOURCE: PCT Int. Appl., 442 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

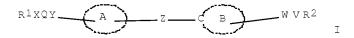
PATENT INFORMATION:

PAT	CENT :	NO.			KINI)	DATE			APPL	ICAT	ION I	. OV		DZ	ATE	
WO	2004	0247)5		A1	_	2004	0325	,	WO 2	003-	JP11	 511		20	0030	909
	W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KR,	KΖ,	LC,	LK,	LR,	LS,
		LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NI,	NO,	NZ,	OM,	PG,
		PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	ΤJ,	TM,	TN,	TR,
		TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW				
	RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑM,	AZ,	BY,
		KG,	KΖ,	MD,	RU,	ТJ,	TM,	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,
		FI,	FR,	GB,	GR,	HU,	ΙE,	ΙΤ,	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,	TR,
		BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	ΤG
JΡ	2004	1237	32		Α		2004	0422	1	JP 2	003-	3164	75		20	0030	909
AU	2003	2620:	23		A1		2004	0430		AU 2	003-	2620:	23		20	0030	909
ΕP	1541	564			A1		2005	0615		EP 2	003-	7953	38		20	0030	909
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙT,	LI,	LU,	NL,	SE,	MC,	PT,

IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
US 2006135578 A1 20060622 US 2005-527426 20050310
PRIORITY APPLN. INFO.: JP 2002-264703 A 20020910
WO 2003-JP11511 W 20030909

OTHER SOURCE(S): MARPAT 140:287404

GΙ



AB The title compds. I [R1 is a group derived from an optionally substituted five-membered heterocycle; X, Y and V are each independently oxygen, sulfur, or the like; Q is a divalent hydrocarbon group having 1 to 20 carbon atoms; A is an aromatic ring which may have one to three addnl. substituents; Z is (CH2)nZ1 or Z1(CH2)n (wherein n is an integer of 0 to 8 and Z1 is oxygen, sulfur, or the like); B is a nitrogenous heterocycle which may have one to three addnl. substituents; W is a bond or a divalent hydrocarbon group having 1 to 20 carbon atoms; and R2 is hydrogen, cyano, PO(OR9)(OR10) (wherein R9 and R10 are each independently hydrogen or optionally substituted hydrocarbyl, or R9 and R10 may be united to form an optionally substituted ring), or the like] are prepared In a binding assay for the human PPAR γ1 receptors, compds. of this invention showed IC50 values of 7.4 nM to 7300 nM. Formulations are given.

IC ICM C07D263-32

ICS C07D413-12; C07D413-14; C07D417-14; C07D417-12; C07D401-14; C07D403-12; C07F007-18; C07F009-6558; A61K031-422; A61K031-4439; A61K031-427; A61K031-4245; A61K031-454; A61K031-5377; A61K031-675; A61K031-695; A61K031-662; A61P003-06; A61P003-04

CC 28-16 (Heterocyclic Compounds (More Than One Hetero Atom))

Section cross-reference(s): 1, 63

REFERENCE COUNT: 65 THERE ARE 65 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 5 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2004:144193 ZCAPLUS Full-text

DOCUMENT NUMBER: 141:925

TITLE: Studies on non-thiazolidinedione antidiabetic agents.

3. Preparation and biological activity of the

metabolites of TAK-559

AUTHOR(S): Imoto, Hiroshi; Matsumoto, Mitsuharu; Odaka,

Biroyuki; Sakamoto, Junichi; Kimura,

Miroyuki; Nonaka, Masami; Kiyota, Yutaka; Momose,

Yu

CORPORATE SOURCE: Pharmaceutical Research Division, Takeda Chemical

Industries, Ltd., Osaka, 532-8686, Japan

SOURCE: Chemical & Pharmaceutical Bulletin (2004), 52(1),

120-124

CODEN: CPBTAL; ISSN: 0009-2363
Pharmaceutical Society of Japan

DOCUMENT TYPE: Journal LANGUAGE: English

PUBLISHER:

OTHER SOURCE(S): CASREACT 141:925

AΒ Preparation and biol. activity of the metabolites of the potent antihyperglycemic and antihyperlipidemic agent, $(E)-4-\{4-[(5-methyl-2-phenyl-2-phe$ 1,3-oxazol-4-yl)methoxy]benzyloxyimino}-4-phenylbutyric acid (TAK-559) (1), were investigated. Metabolites M-I (2), M-II (3), M-III (4) and M-IV (5) were synthesized and their biol. activities were evaluated by in vitro and in vivo expts. Compds. 2-4 activate human peroxisome proliferator-activated receptor gamma one (hPPAR γ 1) and hPPAR α , but their activities are weaker than those of TAK-559 (1). Compound 5 only activates hPPARγ1 weakly. TAK-559 (1) showed potent in vivo plasma glucose and triglyceride lowering activities in Wistar fatty rats after i.p. administration, while its metabolites (2-5) showed comparatively weak activities.

1-10 (Pharmacology)

Section cross-reference(s): 28

THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 16 RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 6 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN 2003:951003 ZCAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER: 140:16723

TITLE: Preparation of 1,2-azole derivatives with hypoglycemic

and hypolipidemic activity

Maekawa, Tsuyoshi; Hara, Ryoma; INVENTOR(S):

Odaka, Hiroyuki; Kimura, Hiroyuki; Mizufune, Hideya; Fukatsu, Kohji

Takeda Chemical Industries, Ltd., Japan; Takeda PATENT ASSIGNEE(S):

Pharmaceutical Company Limited

PCT Int. Appl., 564 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION: DATENT NO

PA:	TENT	NO.			KIN	D	DATE			APPL	ICAT	ION I	.OV		D.	ATE	
WO	2003	0997	93		A8		2003	1229		——— WO 2	003-	JP63	89		2	0030	 522
WO	2003																
	W:	,	•	•	,	,	AU,	,	,	,	,	,	•	•	,		
							DK,										
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KR,	KΖ,	LC,	LK,	LR,	LS,
		LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NΙ,	NO,	NΖ,	OM,	PH
		PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	ΤJ,	TM,	TN,	TR,	TT,	TZ
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	RW:	GH,	GM,	KΕ,	LS,	MW,	${ m MZ}$,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑM,	ΑZ,	BY
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		BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	${ m ML}$,	MR,	ΝE,	SN,	TD,	ΤG
CA	2487	315			A1		2003	1204		CA 2	003-	2487	315		2	0030	522
AU	2003	2411	73		A1		2003	1212		AU 2	003-	2411	73		2	0030	522
JP	2004	2773	97		Α		2004	1007		JP 2	003-	1449	84		2	0030	522
EP	1513	817			A1		2005	0316		EP 2	003-	7305	75		2	0030	522
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PΤ
		ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	HU,	SK	
US	2006	1488	58		A1		2006	0706		US 2	005-	5172	14		2	0050	301
DRIT:	Y APP	LN.	INFO	.:						JP 2	002-	1514	05		A 2	0020	524
										JP 2	002-	2871	61		A 2	0020	930
										JP 2	003-	1674	8		A 2	0030	124
										WO 2	003-	JP63	89	1	w 2	0030	522
ER SO	OURCE	(S):			MARI	PAT	140:	1672	3								

OTHER SOURCE(S): MARPAT 140:16/23 GΙ

1,2-Azole derivs. A-B-Xa-Ya-Xb-Yb-C-Xc-Yc-C(:0)-R (I; e.g. II) wherein ring A AΒ optionally has 1-3 substituents; ring B is a 1,2-azole ring which may further have 1 to 3 substituents; Xa, Xb and Xc are the same or different and each is a bond, -O-, -S- and the like; Ya is a divalent aliphatic hydrocarbon residue having 1-20 C atoms; Yb and Yc are the same or different and each is a bond or a divalent aliphatic hydrocarbon residue having 1-20 C atoms; ring C is a monocyclic aromatic ring which may further have 1 to 3 substituents; and R = -OR4 (R4 is H atom or (un) substituted hydrocarbon group) and the like, or a salt thereof or a prodrug thereof is useful as an agent for the prophylaxis or treatment of diabetes and the like. Hypoglycemic and hypolipidemic actions in mice are tabulated for about 50 examples of I; e.g. a 53 % rate of decrease in blood glucose level in the presence of 0.005 % [2-[3-[3-isopropyl-1-[5-isopropy(trifluoromethyl) - 2-pyridinyl]-1H-pyrazol-4-yl]propoxy]-3-methylphenyl]acetic acid and a 77 % rate of decrease in blood triglyceride level in the presence of 0.005 % 2-methyl-2-[4-[3-methyl-1-[5-(trifluoromethyl)-2-pyridyl]-1Hpyrazol-4- ylmethoxy]phenoxy]propionic acid when the level (glucose or triglyceride) of the non-treated group is taken as 100 %. Plasma antiarteriosclerosis index-enhancing action in mice is tabulated for 34 examples of I, e.g. 25 % for [3-methoxy-2-[3-[3-propy]-1-[5-(trifluoromethy])-2pyridyl]-1H- pyrazol-4-yl]propoxy]phenyl]acetic acid. PPAR γ -RXR α and PPAR δ - $RXR\alpha$ heterodimer ligand activity is tabulated for 59 and 80 examples, resp., of I, e.g. EC50 = 3.8 nM for PPARy-RXR α for [2-[3-[3-cyclohexyl-1-[5-(trifluoromethyl)-2-pyridinyl]-1H-pyrazol-4- yl]propoxy]-3-methylphenyl]acetic acid. Nearly 400 example prepns. of I and 351 example prepns. of intermediates are included. For example, [4-[3-[3-[4-(trifluoromethyl)phenyl]-5-isoxazolyl]propoxy]phenyl]acetic acid was obtained in 25 % yield from a mixture of 3-[3-[4- (trifluoromethyl)phenyl]-5isoxazolyl]-1-Pr methanesulfonate, NaI, Me 2-(4-hydroxyphenyl)acetate, K2CO3 and DMF; details of the preparation of the mesylate are also given. ICM C07D231-12 IC C07D261-08; C07D401-04; C07D413-12; A61K031-4155; A61K031-415;

A61K031-42; A61K031-422; A61K031-4439; C07D231-14; C07D231-20; C07D231-22; C07D401-14; C07D403-04; C07D403-14

28-8 (Heterocyclic Compounds (More Than One Hetero Atom))

Section cross-reference(s): 1, 63

THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 19 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

10/517214 L88 ANSWER 7 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2003:270930 ZCAPLUS Full-text DOCUMENT NUMBER: 139:46234 TITLE: Activation of human PPAR subtypes by Pioglitazone AUTHOR(S): Kimura, Hiroyuki; Sakamoto, Junichi; Moriyama, Shinji; Odaka, Hiroyuki; Momose, Yu; Sugiyama, Yasuo; Ikeda, Hitoshi; Sawada, Hidekazu CORPORATE SOURCE: Discovery Research Laboratories IV, Pharmaceutical Discovery Research Division, Takeda Chemical Industries, Ltd., Yodogawa-ku, Osaka, 532-8686, Japan SOURCE: Medical Science Symposia Series (2002), 18 (Peroxisome Proliferator Activated Receptors), 41-47 CODEN: MSSYEI; ISSN: 0928-9550 Kluwer Academic Publishers PUBLISHER: DOCUMENT TYPE: Journal; General Review LANGUAGE: English AΒ A review. Pioglitazone activates both human peroxisome proliferator activated receptor γ (hPPAR γ) and hPPAR α . Pioglitazone improves insulin sensitivity in patients with type 2 diabetes, and significantly decreased mean triglycerides levels and increased high-d. lipoprotein-cholesterol levels in both monotherapy and in combination with sulfonylureas, metformin or insulin. The good effects for lipid profile of Pioglitazone are partly mediated by PPARlpha. CC 1-0 (Pharmacology) Section cross-reference(s): 2, 14 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 15 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L88 ANSWER 8 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN 2003:5954 ZCAPLUS Full-text ACCESSION NUMBER: DOCUMENT NUMBER: 138:89798 TITLE: Preparation of 4-(phenoxymethyl)-5-methyloxazole derivatives as antidiabetic agents INVENTOR(S): Momose, Yu: Maekawa, Tsuvoshi: Odaka, Hiroyuki; Kimura, Hiroyuki PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Japan SOURCE: PCT Int. Appl., 99 pp. CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: Japanese FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. A1 20030103 WO 2002-JP6107 WO 2003000685 20020619 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,

> CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG

AU 2002-315787 20020619

JP 2002-178851 20020619 JP 2001-186952 A 20010620 WO 2002-JP6107 W 20020619

20020619

JP 2002-178851

OTHER SOURCE(S): MARPAT 138:89798

JP 2003073377

PRIORITY APPLN. INFO.:

AU 2002315787 A1 20030108

A

20030312

GΙ

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AΒ The title compds. I [wherein R1 = (un) substituted (hetero) hydrocarbonyl; X and Y = independently a bond, O, S, CO, CS, SO, SO2, CR3OR4, NR5, CONR6, or NR6CO; R3 and R6 = independently H or (un)substituted hydrocarbonyl; R4 = H or protecting group of OH; R5 = H, (un)substituted hydrocarbonyl, or protecting group of amino; Q and W = independently (CH2)m; m = 1-20; ring A =(un) substituted aryl; n = 1-8; ring B = (un) substituted 5-membered ring containing N; V = a bond, O, S, SO, SO2, NR7, or NR7CO; R7 = H or (un) substituted hydrocarbonyl; R2 = PO(OR8)(OR9), COR10, (un) substituted hydrocarbonyl, or heteroaryl; R8 and R9 = independently H or (un)substituted hydrocarbonyl; or R8 and R9 together form (un)substituted ring; R10 = H or (un) substituted hydrocarbonyl; with provisos] and salts or prodrugs thereof are prepared as antidiabetic agents. For example, the acid II (prepn given) was treated with iso-Bu chlorocarbonate in THF in the presence of 4methylmorpholine, followed by the addition of THF solution of H2NNH2•H2O. The above product was then reacted with tri-Me orthobutyrate in 1.4-dioxane in the presence of methanesulfonic acid to afford the target compd III (70%). III showed IC50 of 0.034 μM and 0.15 μM against peroxisome proliferator-activated receptors (PPAR) γ and PPAR γ -RXR α , resp. A capsule formulation containing III as an active ingredient was also described.

IC ICM C07D413-12 ICS C07D413-14; C07D417-14; A61K031-422; A61K031-427; A61K031-4439;

CC 28-6 (Heterocyclic Compounds (More Than One Hetero Atom))

Section cross-reference(s): 1, 63

REFERENCE COUNT: 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 9 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2003:5768 ZCAPLUS <u>Full-text</u>

A61P003-04; A61P003-06; A61P003-10

DOCUMENT NUMBER: 138:66691

TITLE: Function regulator for retinoid relative receptor

INVENTOR(S): Maekawa, Tsuyoshi; Kunitomo, Jun; Odaka, Hiroyuki; Kimura, Hiroyuki

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Japan

SOURCE: PCT Int. Appl., 114 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

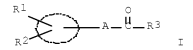
PATENT INFORMATION:

PAT	TENT I	. O <i>V</i>			KIN	D	DATE			APPL	ICAT	ION 1	NO.		D	ATE	
						_											
WO	2003	0002	49		A1		2003	0103	1	WO 2	002-	JP63	49		2	0020	625
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KR,	KΖ,	LC,	LK,	LR,	LS,
		LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,	PL,
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		UG,	US,	UZ,	VN,	YU,	ZA,	ZM,	ZW								
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AT,	BE,	CH,
		CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,
		BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	ΤG

AU 2002315885 A1 20030108 AU 2002-315885 20020625 JP 2003081832 A 20030319 JP 2002-184633 20020625 EP 1405636 A1 20040407 EP 2002-741287 20020625 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR US 2003-481033 US 2004157881 Α1 20040812 20031216 US 7223791 В2 20070529 JP 2001-192601 A 20010626 WO 2002-JP6349 W 20020625 PRIORITY APPLN. INFO.:

OTHER SOURCE(S): MARPAT 138:66691

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AB A function regulator for retinoid relative receptors (excluding retinoic acid receptors) which contains a compound represented by the general formula I [one of R1 and R2 = monocyclic aromatic hydrocarbon group (substituted) or monocyclic aromatic heterocyclic group containing one heteroatom and the other represents hydrogen (substituted), etc.; B = 5- or 6-membered heterocycle (excluding 1,3-azole); A = aromatic hydrocarbon group (substituted) or aromatic heterocyclic group (substituted); and R3 = hydrogen, etc.] or a salt thereof. The regulator is useful as a preventive/remedy for diabetes, hyperlipidemia, impaired glucose tolerance, etc.

IC ICM A61K031-341

ICS A61K031-381; A61K031-40; A61K031-4196; A61K031-42; A61K031-4245; A61K031-433; A61K031-4418; A61K031-625; A61K045-00; A61P003-00; A61P003-04; A61P003-06; A61P003-10; A61P005-50; A61P043-00; C07D207-337; C07D307-54; C07D213-61; C07D249-08

CC 1-10 (Pharmacology)

Section cross-reference(s): 28, 63

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 10 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:754366 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 137:279197

TITLE: Preparation of five-membered heterocyclic alkanoic

acid derivatives as remedies for diabetes and $% \left(1\right) =\left(1\right) +\left(1$

hyperlipidemia

INVENTOR(S): Momose, Yu; Maekawa, Tsuyoshi; Imoto,

Hiroshi; Odaka, Hiroyuki; Kimura,

Hiroyuki

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Japan

SOURCE: PCT Int. Appl., 165 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002076959	A1	20021003	WO 2002-JP2741	20020322

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W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
            GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS,
            LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL,
            PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA,
            UG, US, UZ, VN, YU, ZA, ZM, ZW
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,
            CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
            BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                               20021008 AU 2002-239023
    AU 2002239023
                         A1
                                                                  20020322
    JP 2002348281
                               20021204
                                          JP 2002-81621
                         Α
                                                                  20020322
                                        EP 2002-705433
    EP 1394154
                               20040303
                         Α1
                                                                  20020322
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
    US 2004063775
                        A1
                               20040401
                                          US 2003-472159
                                                                  20030922
    US 7241785
                        В2
                               20070710
PRIORITY APPLN. INFO.:
                                           JP 2001-85572
                                                             A 20010323
                                                             W 20020322
                                           WO 2002-JP2741
                       MARPAT 137:279197
OTHER SOURCE(S):
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 R^1XQY A Z C B $W(C=0)R^2$

The title compds. I [R1 represents an optionally substituted five-membered heterocyclic group; X represents a bond, etc.; Q represents a C1-20 divalent hydrocarbon group; Y represents a bond, etc.; ring A represents an aromatic ring optionally having one to three substituents; Z represents (CH2)nZ1 (n is an integer of 0 to 8 and Z1 represents a bond, etc.), etc.; ring B represents a five-membered heterocycle optionally having one to three substituents; W represents a C1-20 divalent saturated hydrocarbon group; and R2 represents OH, etc.] are prepared A process for preparing I is disclosed. Compds. of this invention at 0.01% in feed given to diabetic mice for 4 days caused 43% to 42% decrease of blood sugar. Formulations are given.

IC ICM C07D263-32 ICS C07D263-34; C07D413-12; C07D413-14; C07D417-12; A61K031-421; A61K031-422; A61K031-427; A61K031-4439; A61K031-4709; A61K031-5377; A61P003-06; A61P003-10; A61P043-00

CC 28-10 (Heterocyclic Compounds (More Than One Hetero Atom))

Section cross-reference(s): 1, 63

REFERENCE COUNT: 88 THERE ARE 88 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 11 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:521714 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 137:109278

INVENTOR(S):

TITLE: Preparation of alkanoic acid derivatives as preventives and/or remedies for diabetes,

hyperlipidemia, impaired glucose tolerance, and

retinoid-related receptor regulators
Momose, Yu; Maekawa, Tsuyoshi; Takakura,

Nobuyuki; Odaka, Hiroyuki; Kimura,

Hiroyuki; Ito, Tatsuya

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Japan

SOURCE: PCT Int. Appl., 235 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

GI

PA.	TENT :	NO.			KIN	D	DATE			APPL	ICAT	ION :	NO.		Γ	ATE	
WO	2002	 0535	47		A1	_	2002	0711		WO 2	001-	 JP11	 611		2	20011	228
	W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	AZ,	ΒA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
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		LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NΖ,	OM,	PH,	PL,
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	RW:						MZ,							•			•
		CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	ΙT,	LU,	MC,	NL,	PT,	SE,	TR,
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	2433				A1		2002									20011	228
	2002															20011	
JP	2002	2654	57		А		2002	0918		JP 2	001-	4020	99		2	20011	228
EP	1357	115			A1		2003	1029		EP 2	001-	2725	44		2	20011	228
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,
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US	2004									US 2	003-	4659	38		2	20030	626
US	7238	716			В2		2007	0703									
PRIORIT	Y APP	LN.	INFO	.:							000-						
										WO 2	001-	JP11	611		W 2	20011	228
OTHER SO	DURCE	(S):			MAR	PAT	137:	1092	78								

R1 - X-O-Y - A Z - R II-W-CO-R3

AB Alkanoic acid derivs. represented by the general formula (I) or salts thereof [wherein R1 = optionally substituted five-membered aromatic heterocyclic group; X = a bond, O, S, CO, C(:S), CR4(OR6), NR6 (wherein R4 = H, optionally substituted hydrocarbyl; R5 = H, hydroxy-protecting group; R6 = H, optionally hydrocarbyl, amino-protecting group); Q = C1-20 divalent hydrocarbon group; Y = bond, O, S, S(:0), SO2, NR7, CONR7, NR7CO, (wherein R7 = H, optionally substituted hydrocarbon group, amino-protecting group); ; ring A = an aromatic ring which may have one to three substituents; Z = (CH2)n-Z1 (wherein n = aninteger of 1 to 8; Z1 = O, S, SO, SO2, NR16; wherein R16 = H, optionally substituted hydrocarbon group); ring B = an optionally mono- to trisubstituted pyridine, benzene, or naphthalene ring; U = a bond, O, S, SOP, SO2; W = C1-20 divalent hydrocarbon group; R3; R3 = OH, optionally substituted hydrocarbyloxy, NR9R10 (wherein R9, R10 = H, optionally substituted hydrocarbyl, heterocyclyl, or acyl; or R9 and R10 are linked to each other to form a ring); with the proviso that when B is an optionally mono- to trisubstituted benzene ring, U is a bond] are prepared Also disclosed are preventives and/or remedies for diabetes, hyperlipidemia, and impaired glucose tolerance, retinoid-related receptor regulators, ligands for peroxisomeproliferator response receptor and retinoid X receptor, insulin resistance improvers containing the compds. I or salts or prodrugs thereof. Thus, a 40%

toluene solution (1.74 g) of di-Et azodicarboxylate was added dropwise to a mixture of 3-(5-methyl-2-phenyl-4-oxazolylmethoxy)-5- isoxazolylmethanol 0.859, Me 2-(2-hydroxyphenyl)acetate 0.499, Ph3P 0.944, and 15 mL THF at room temperature and stirred for 15 h to give Me 2-[2-[3-(5-methyl-2-phenyl-4-oxazolylmethoxy)-5- isoxazolylmethoxy]phenyl]acetate as an oil which was dissolved in MeOH/THF (1/1, 20 mL), treated with 10 mL 1 N aqueous NaOH, stirred at room temperature for 15 h, and acidified with 1 N aqueous HCl to give 52% 2-[2-[3-(5-methyl-2-phenyl-4-oxazolylmethoxy)-5- isoxazolylmethoxy]phenyl]acetic acid (II). When a feed containing 0.005% II was fed freely to type II diabetic mice for 4 days, the blood sugar and lipid level was lowered by 54 and 96%, resp. A capsule and a tablet formulation containing 2-[2-ethoxy-5-[4-[(5-methyl-2-phenyl-4-

oxazolyl)methoxy]benzyloxy]phenyl]acetic acid Me ester were prepared

IC ICM C07D263-32

CC 28-10 (Heterocyclic Compounds (More Than One Hetero Atom))

Section cross-reference(s): 1, 63

REFERENCE COUNT: 11 THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 12 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:391693 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 136:401786

TITLE: Preparation of isoxazole derivatives for prevention

and treatment of diabetes

INVENTOR(S): Momose, Yu; Maekawa, Tsuyoshi; Asakawa,

Tomoko; Sakai, Nozomu

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Japan

SOURCE: PCT Int. Appl., 270 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	CENT	NO.			KINI)	DATE			APPL	ICAT	ION 1	.OV		D.	ATE	
WO	2002	0404	 58		A1		2002	0523	,	WO 2	001-	JP10	001		2	0011	116 <
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		LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,	PL,
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		US,	UΖ,	VN,	YU,	ZA,	ZM,	ZW									
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							FR,										
		BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG
CA	2429	426			A1		2002	0523	1	CA 2	001-	2429	426		2	0011	116 <
AU	2002	0152	18		A5		2002	0527		AU 2	002-	1521	8		2	0011	116
JP	2002	2121	71		А		2002	0731		JP 2	001-	3524	66		2	0011	116
EP	1340	749			A1		2003	0903		EP 2	001-	9838	8 0		2	0011	116
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR						
US	2004	0489	08		A1		2004	0311		US 2	003-	4166	58		2	0030	514
US	7022	725			В2		2006	0404									
US	2006	0846	90		A1		2006	0420		US 2	005-	2950	58		2	0051	206
PRIORITY	APP	LN.	INFO	.:						JP 2	000-	3508	69	1	A 2	0001	117

WO 2001-JP10001 W 20011116 US 2003-416658 A3 20030514

OTHER SOURCE(S): MARPAT 136:401786

GΙ

$$\mathbb{R}^{1} \longrightarrow \mathbb{R}^{2}$$

$$\mathbb{R}^{2}$$

$$\mathbb{R}^{2}$$

Described are preventives or remedies for diabetes containing compds. of the AΒ general formula (I) or their salts or prodrugs thereof [wherein one of R1 and R2 is hydrogen or a substituent and the other is an optionally substituted cyclic group; W is a free valency or a divalent aliphatic hydrocarbon group; and Y is a group represented by the general formula OR3 (wherein R3 is hydrogen, optionally substituted hydrocarbyl, an optionally substituted heterocyclic group, or optionally substituted acyl) or carboxyl which may be converted into an ester or an amide]. These compds. have excellent insulin secretion-promoting and blood sugar-decreasing effects and low toxicity and are useful as drugs, particularly preventive and therapeutic agents for diabetes and diabetic complication. Thus, reduction of 3-[5-(3,4dichlorophenyl)-4-isoxazolyl]propionic acid Me ester (preparation given) by diisobutylaluminum hydride in hexane/THF at room temperature for 1 h gave 97% 3-[5-(3,4-chlorophenyl)-4-isoxazolyl] propanol (II). II at 30 mg/kg p.o. was administered to rats and after 60 min, the rats were fed with glucose at 2q/kg p.o. After 30 min, the blood sample was taken and the blood sugar level measured was 75% of the control. A capsule and tablet formulation containing II were formulated.

IC ICM C07D261-08
ICS C07D417-12; C07D413-12; C07D413-06; C07D413-14; C07D413-04;
A61K031-42; A61K031-675; A61K031-427; A61K031-4709; A61K031-496;
A61K031-454

CC 28-18 (Heterocyclic Compounds (More Than One Hetero Atom))
Section cross-reference(s): 1, 63

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RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of isoxazole derivs. having insulin secretion promoting and blood sugar decreasing effects for prevention and treatment of diabetes and diabetes complication)

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RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(uses)

ΙT

(preparation of isoxazole derivs. having insulin secretion promoting and

blood sugar decreasing effects for prevention and treatment of diabetes and diabetes complication)

REFERENCE COUNT: 11 THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 13 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:149264 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 136:340623

TITLE: Novel 5-Substituted 2,4-Thiazolidinedione and 2,4-Oxazolidinedione Derivatives as Insulin

2,4-Oxazolidinedione Derivatives as Insulin Sensitizers with Antidiabetic Activities Momose, Yu; Maekawa, Tsuyoshi; Yamano,

AUTHOR(S): Momose, Yu; Maekawa, Tsuyoshi; Yamano, Tohru; Kawada, Mitsuru; Odaka, Hiroyuki;

Ikeda, Hitoshi; Sohda, Takashi

CORPORATE SOURCE: Medicinal Chemistry Research Laboratories II,

Pharmacology Research Laboratories II, and Strategic Research Planning, Pharmaceutical Research Division, Takeda Chemical Industries Ltd., Yodogawaku, Osaka,

532-8686, Japan

SOURCE: Journal of Medicinal Chemistry (2002), 45(7),

1518-1534

CODEN: JMCMAR; ISSN: 0022-2623

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 136:340623

GΙ

AΒ $5-(\omega-Azolylalkoxyphenylalkyl)-2,4-thiazolidinones$ and -2,4-oxazolidinones such as furylmethyloxazolylmethoxymethoxyphenylpropyl oxazolidinedione I were prepared as potential antidiabetic and antihyperlipidemic agents. Many of the 2,4-thiazolidinediones and 2,4-oxazolidinones showed potent glucose- and lipid-lowering activities. The antidiabetic activities of the 2,4oxazolidinediones were superior to those of the 2,4-thiazolidinediones. Both enantiomers of I, one of the most interesting compds. in terms of activity, were synthesized by using an asym. O-acetylation of the corresponding α hydroxyvalerate with immobilized lipase, followed by cyclization of the oxazolidinedione ring. The (R)-(+)-enantiomer of I showed more potent glucoselowering activity [ED25 = 0.561 mg/kg/d] than either the (S)-(-)-enantiomer (ED25 > 1.5 mg/kg/d) or pioglitazone (ED25 = 6 mg/kg/d) in KKAy mice. (+)(R)-I also exhibited a 10-fold more potent antidiabetic activity (ED25 = 0.05mg/kg/d) than pioglitazone (ED25 = 0.5 mg/kg/d) in Wistar fatty rats. The antidiabetic effects of I are related to its activity as a potent agonist for peroxisome proliferator-activated receptor γ (PPAR- γ) (EC50 = 8.87 nM). The

Ι

crystal structures of intermediates in the synthesis of nonracemic thiazolidinediones were determined by X-ray crystallog.

C 28-7 (Heterocyclic Compounds (More Than One Hetero Atom))

Section cross-reference(s): 1, 75

REFERENCE COUNT: 29 THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 14 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:19837 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 136:350405

TITLE: Novel 5-substituted-1H-tetrazole derivatives as potent

glucose and lipid lowering agents

AUTHOR(S): Momose, Yu.; Maekawa, Tsuyoshi; Odaka, Miroyuki; Ikeda, Hitoshi; Sohda, Takashi

COPPORATE SOURCE: Madicinal Chamistry Possarch Laboratories II

CORPORATE SOURCE: Medicinal Chemistry Research Laboratories II, Takeda

Chemical Industries, Ltd., Chuo-ku. Osaka, 540-8645,

Japan

SOURCE: Chemical & Pharmaceutical Bulletin (2002), 50(1),

100-111

CODEN: CPBTAL; ISSN: 0009-2363
Pharmaceutical Society of Japan

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 136:350405

GΙ

PUBLISHER:

AB A series of 5-(4-alkoxyphenylalkyl)-1H-tetrazole derivs. containing an oxazole-based group at the alkoxy moiety was prepared; the antidiabetic and antihyperlipidemic effects of members of the series were evaluated in two genetically obese and diabetic animal models. The tetrazole compds. were prepared using the cycloaddns. of azides with the corresponding nitriles. Many of the 5-(4-alkoxyphenylalkyl)-1H-tetrazoles showed potent glucose and lipid lowering activities in KKAy mice. Methylphenyloxazolylmethoxypy ridylpropyltetrazole I had potent glucose lowering activity (ED25 = 0.0839 mg·kg-1·d-1), being 72 times more active than pioglitazone hydrochloride (ED25 = 6.0 mg·kg·d-1); in addition, I also exhibited strong antihyperlipidemic activity (ED25 = 0.0277 mg·kg-1·d-1) in Wistar fatty rats. The antidiabetic activity of I is likely related to its potent agonistic activity for peroxisome proliferator-activated receptor γ (PPAR γ) (EC50 = 6.75 nM).

CC 1-10 (Pharmacology)

Section cross-reference(s): 28

REFERENCE COUNT: 33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 15 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:396864 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 135:19632

TITLE: Preparation of pyrazolyl- and pyrrolylalkanoic acid

derivatives with hypoglycemic and hypolipidemic

activity

INVENTOR(S): Momose, Yu; Maekawa, Tsuyoshi; Odaka,

Hiroyuki; Kimura, Hiroyuki

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Japan

SOURCE: PCT Int. Appl., 375 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

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	TENT						DATE			APPL	ICAT	ION	NO.		D	ATE	
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$$X1 - R2$$

 $R^1 - X - (CH_2)_m - Y - A - (CH_2)_n - B - W - CO - R^3$

I

$$\begin{array}{c} \text{Me} \\ \text{CH}_2 = \text{O} \\ \end{array}$$

AΒ Title compds. (I) [wherein R1 = (un)substituted hydrocarbon or heterocycle; X = bond, O, S, CO, CS, CR4(OR5), or NR6; R4 and R6 = independently H or (un) substituted hydrocarbon; R5 = H or hydroxyl protective group; m = 0-3; Y =O, S, SO, SO2, NR7, CONR7, or NR7CO; R7 = H or (un)substituted hydrocarbon; A = (un)substituted aromatic ring; n = 1-8; B = (un)substituted N-containing 5membered heterocycle; X1 = bond, O, S, SO, SO2, OSO2, or NR16; R16 = H or (un) substituted hydrocarbon; R2 = H or (un) substituted hydrocarbon or heterocycle; W = bond or hydrocarbon; R3 = OR8 or NR9R10; R8 = H or (un) substituted hydrocarbon; R9 and R10 = independently H or (un) substituted hydrocarbon or heterocycle; or R9 and R10 together with the N to which they are attached may form a ring] were prepared as retinoid-related receptor function regulating agents or insulin resistance improving agents. For example, Et 3-[1-(4-hydroxybenzyl)-4- phenyl-3-pyrrolyl]propionate and 4chloromethyl-5-methyl-2-(2-thienyl) oxazole were coupled in the presence of K2CO3 in DMF and treated with HCl to give II (77%). At a concentration of 0.001%, II reduced hypoglycemic and hypolipidemic action by 48% and 70%, resp., lowered total cholesterol by 16%, and increased the plasma antiarteriosclerosis index by 12% compared to non-treatment groups of mice. In addition, II showed potent PPARy-RXR α heterodimer ligand activity with EC50 of 1.5 nM. I are useful for the prevention or treatment of diabetes mellitus, hyperlipidemia, impaired glucose tolerance, inflammatory diseases, and arteriosclerosis.

IC ICM C07D409-12

ICS C07D413-12; C07D401-14; C07D405-12; C07D231-12; C07D401-12; C07D417-14; C07D409-14; A61K031-501; A61P003-00

CC 28-8 (Heterocyclic Compounds (More Than One Hetero Atom))
 Section cross-reference(s): 1

3256-88-0P, 2-Methyl-5-phenylpyridine 4634-09-7P ΙT 5229-40-3P 111770-91-3P 116140-28-4P 146775-28-2P, 80457-61-0P 2-Chloromethyl-5-phenylpyridine 162614-73-5P 177275-37-5P 177976-31-7P, 3-Chloromethyl-5-phenylpyridine 187392-96-7P 194546-13-9P 197847-89-5P 339269-10-2P 339269-11-3P 342023-31-8P 342023-32-9P 342023-34-1P 342023-36-3P 342023-37-4P 342023-39-6P 342023-41-0P 342023-43-2P 342023-44-3P 342023-46-5P 342023-48-7P 342023-49-8P 342023-52-3P 342023-54-5P 342023-56-7P 342023-58-9P 342023-59-0P 342023-61-4P 342023-63-6P 342023-65-8P 342023-67-0P 342023-69-2P 342023-68-1P 342023-70-5P 342023-72-7P 342023-73-8P 342023-75-0P 342023-76-1P 342023-78-3P 342023-79-4P 342023-80-7P 342023-81-8P 342023-82-9P 342023-83-0P 342023-84-1P 342023-85-2P 342023-86-3P 342023-87-4P 342023-88-5P 342023-90-9P 342023-91-0P 342023-92-1P 342023-98-7P 342023-99-8P 342024-01-5P 342024-00-4P 342024-02-6P 342024-04-8P 342024-06-0P 342024-07-1P 342024-08-2P 342024-09-3P

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                          342024-79-7P 342024-82-2P 342024-84-4P
342024-85-5P 342024-86-6P
                          342024-88-8P 342024-89-9P
                                                     342024-90-2P
342024-91-3P 342024-92-4P 342024-93-5P 342024-94-6P 342024-95-7P
342024-96-8P 342024-97-9P 342024-98-0P 342024-99-1P 342025-01-8P
342025-02-9P 342025-04-1P 342025-05-2P 342025-06-3P 342025-07-4P
342025-08-5P 342025-10-9P 342025-11-0P 342025-12-1P 342027-87-6P
342028-02-8P
```

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate; preparation of pyrrolyl- and pyrazolylalkanoic acid derivs. as retinoid X receptor and PPAR receptor modulators)

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 16 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:359973 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 134:353301

Preparation of alkoxyiminoalkanoic acid derivatives TITLE:

having blood sugar and lipid lowering effect Momose, Yu; Imoto, Hiroshi; Odaka, Miroyuki;

Kimura, Hiroyuki

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Japan

PCT Int. Appl., 92 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION: DAMENIE NO

INVENTOR(S):

PA	TENT :	NO.			KIN	D	DATE			APPL	ICAT	ION I	NO.		D.	ATE	
WO	2001	 0345	 79		A1	_	2001	0517	,	WO 2	000-	 JP78	 78		2	0001	109
	W:	ΑE,	AG,	AL,	AM,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CN,	CR,	CU,
		CZ,	DM,	DZ,	EE,	GD,	GE,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KG,	KR,	KΖ,
		LC,	LK,	LR,	LT,	LV,	MA,	MD,	MG,	MK,	MN,	MX,	NO,	NZ,	PL,	RO,	RU,
		SG,	SI,	SK,	ΤJ,	TM,	TR,	TT,	UA,	US,	UZ,	VN,	YU,	ZA,	AM,	AZ,	BY,
		KG,	KΖ,	MD,	RU,	ΤJ,	TM										
	RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,
		ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,		
		ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,	ΤG		
CA	2390	928			A1		2001	0517	1	CA 2	000-	2390	928		2	0001	109
AU	2001	0130	32		A5		2001	0606		AU 2	001-	1303	2		2	0001	109
JP	2001	1999	71		Α		2001	0724		JP 2	000-	3474	63		2	0001	109
EP	1229	026			A1		2002	0807		EP 2	000-	9748.	58		2	0001	109
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR						
PRIORIT	Y APP	LN.	INFO	.:					1	JP 1	999-	3203	18	Ž	A 1	9991	110
									,	WO 2	000-	JP78	78	Ī	W 2	0001	109
OTHER S	OURCE	(S):			MAR	PAT	134:	35330	01								

GΙ

AΒ Compds. of general formula (I) or salts thereof [wherein R1 = an optionally substituted hydrocarbon or heterocyclyl; X is a free valency, O, S, CO, CS, CR6(OR7), NR8 (wherein R6, R8 = H, optionally substituted hydrocarbyl; R7 = H, HO-protective group); Y = O, S, SO, SO2, NR8, CONR8, NR8CO (wherein R8 = same as above); ring A = a heterocycle or hydrocarbon ring optionally having 1-3 substituents; p = 1-8; R2 = hydrogen, optionally substituted hydrocarbyl or heterocyclyl; q = 0-6; m = 0,1; R3 = OH, optionally substituted hydrocarbyloxy or NH2; R4, R5 = H, optionally substituted hydrocarbyl; or R4 and R2 are linked together to form a ring; with the provisos that when A is optionally substituted indole, Y is not oxygen or sulfur, that when Y is oxygen, sulfur, -SO-, -SO2-, or -NR8-, A is not an optionally substituted benzene ring, and that when Y is oxygen and A is an optionally substituted, 4-pyrone, 4pyridone, or pyridine N-oxide ring, R2 is not a thiazolyl or thiadiazolyl group substituted with optionally protected amino] are prepared These compds. are ligand for peroxisome proliferator-activated receptor (PPARV) and retinoid-related receptors, in particular retinoid X receptors and useful as preventive or therapeutic agents for diabetes, hyperlipidemia, or glucose intolerance and as insulin resistance improvers. Thus, NaH was gradually added to a solution of 5-chloromethyl-2-(5-methyl-2-phenyl-4oxazolylmethoxy)pyridine and (E)-4-(hydroxyimino)-4-phenylbutanoic acid Me ester in DMF at 0° and stirred at room temperature for 1.5 h to give 87% (E)-4-[6-(5-methyl-2-phenyl-4-oxazolylmethoxy)-3-pyridylmethoxyimino]-4phenylbutanoic acid Me ester which was saponified with LiOH in aqueous methanol and acidified with 1 N HCl to give 87% (E)-4-[6-(5-methyl-2-phenyl-4oxazolylmethoxy)-3-pyridylmethoxyimino]-4-phenylbutanoic acid (II). KKAY mice (obesity and diabetes type II model), who were fed with a powder feed containing 0.01% II for 4 days, lowered blood sugar and triglyceride level by 54 and 90%, resp. A capsule and tablet formulation containing II were prepared

IC ICM C07D263-32

ICS C07D413-12; A61K031-421; A61K031-4439; C07C251-54; A61K031-195; A61K031-235; A61P003-10; A61P003-06

CC 28-6 (Heterocyclic Compounds (More Than One Hetero Atom))

Section cross-reference(s): 1

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 17 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:359842 ZCAPLUS Full-text

DOCUMENT NUMBER: 134:361377

TITLE: Body weight gain inhibitors

INVENTOR(S): Sugiyama, Yasuo; Odaka, Miroyuki;

Kimura, Hiroyuki

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Japan

SOURCE: PCT Int. Appl., 67 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA:	TENT	NO.			KIN	D	DATE			APPL	ICAT	ION 1	NO.		Ι	DATE	
WO	2001	0342	00		A1	_	2001	0517		 WO 2	000-	 JP78	 79		2	20001	109
	W:	ΑE,	AG,	AL,	AM,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CN,	CR,	CU,
		CZ,	DM,	DZ,	EE,	GD,	GE,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KG,	KR,	KΖ,
		LC,	LK,	LR,	LT,	LV,	MA,	MD,	MG,	MK,	MN,	MX,	NO,	NZ,	PL,	RO,	RU,
		SG,	SI,	SK,	ΤJ,	TM,	TR,	TT,	UA,	US,	UZ,	VN,	YU,	ZA			
	RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,
		DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,	TR,	BF,
		ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,	ΤG		
CA	2390	932			A1		2001	0517		CA 2	000-	2390	932		2	20001	109
AU	2001	0130	33		Α5		2001	0606		AU 2	001 -	1303.	3		2	20001	109
JP	2001	1998	87		Α		2001	0724		JP 2	000-	3474	64		2	20001	109
HU	2002	0383	7		A2		2003	0328		HU 2	002-	3837			2	20001	109
EP	1304	121			A1		2003	0423		EP 2	000 -	9748	59		2	20001	109
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR						
ИО	2002	0022	14		Α		2002	0709		NO 2	002-	2214			2	20020	508
US	2005		A1		2005	1027		US 2	005-	1683	57		2	20050	629		
IORIT:	Y APP	LN.	INFO	.:						JP 1	999-	3203	19	i	A 1	9991	110
										WO 2	000-	JP78	79	Ţ	W 2	20001	109
										US 2	002-	1297	04]	B1 2	20020	509
HER SO	DURCE	(S):			MAR	PAT	134:	3613	77								

OTHER SOURCE(S):

GΙ

IC

$$R^{1}X(CH_{2})_{n}Y$$
 (CH₂) ponc (CH₂) q (C) mCOR³

Body weight gain inhibitors comprises PPARy agonist-like substances, which AΒ contain PPAR δ agonist-like substances such as compds. represented by general formula (I) wherein R1 represents optionally substituted hydrocarbyl, etc.; X represents a bond, etc.; Y represents oxygen, etc.; the ring A represents a heterocycle, etc.; R2 represents hydrogen, etc.; R3 represents-OR9, etc.; and R4 and R5 represent each hydrogen, etc., are useful in treating diabetes, etc.

ICS C07D263-32; A61K031-421; A61K031-195; A61K031-235; A61P003-10; A61P003-04

CC 1-10 (Pharmacology)

ICM A61K045-00

Section cross-reference(s): 28, 63

REFERENCE COUNT: THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 18 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN 2000:832682 ZCAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER: 134:157419

Activation of Human Peroxisome Proliferator-Activated TITLE:

Receptor (PPAR) Subtypes by Pioglitazone

AUTHOR(S): Sakamoto, Junichi; Kimura, Hiroyuki;

Moriyama, Shinji; Odaka, Hiroyuki; Momose,

Yu; Suqiyama, Yasuo; Sawada, Hidekazu

CORPORATE SOURCE: Discovery Research Laboratories IV, Pharmaceutical

Discovery Research Division, Takeda Chemical

Industries, Ltd., Osaka, Japan

SOURCE: Biochemical and Biophysical Research Communications

(2000), 278(3), 704-711

CODEN: BBRCA9; ISSN: 0006-291X

PUBLISHER: Academic Press

DOCUMENT TYPE: Journal LANGUAGE: English

AB Pioglitazone, a thiazolidinedione (TZD) derivative, is an antidiabetic agent that improves hyperglycemia and hyperlipidemia in obese and diabetic animals via a reduction in hepatic and peripheral insulin resistance. The TZDs including pioglitazone have been identified as high affinity ligands for peroxisome proliferator-activated receptor (PPAR) γ . The selectivity of pioglitazone for the human PPAR subtypes has not been reported, thus, we investigated the effect of pioglitazone on the human PPAR subtypes. Transient transactivation assay showed that pioglitazone is a selective hPPAR γ 1 activator and a weak hPPAR α activator. Binding assay indicated that the transactivation of hPPAR γ 1 or hPPAR α by pioglitazone is due to direct binding of pioglitazone to each subtype. Furthermore, pioglitazone significantly increased the apoA-I secretion from the human hepatoma cell line HepG2. (c) 2000 Academic Press.

CC 1-10 (Pharmacology)

REFERENCE COUNT: 29 THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 19 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2000:34864 ZCAPLUS Full-text

DOCUMENT NUMBER: 132:93338

TITLE: Preparation of heterocyclic compounds as retinoid-associated receptor regulators

INVENTOR(S): Sugiyama, Yasuo; Momose, Yu; Kimura, Hiroyuki

; Sakamoto, Junichi; Odaka, Miroyuki

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Japan

SOURCE: PCT Int. Appl., 122 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	CENT	NO.			KIN	D	DATE		-	APPL	ICAT	ION :	NO.		D.	ATE	
WO	2000	 0016	 79		 A1	_	2000	0113	;	 WO 1	 999-	 JP35	 20		1	9990	630
	W:	ΑE,	AL,	ΑM,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CN,	CU,	CZ,	EE,	GD,
		GE,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KG,	KR,	KΖ,	LC,	LK,	LR,	LT,	LV,
		MD,	MG,	MK,	MN,	MX,	NO,	NZ,	PL,	RO,	RU,	SG,	SI,	SK,	SL,	ТJ,	TM,
		TR,	TT,	UA,	US,	UZ,	VN,	YU,	ZA								
	RW:	GH,	GM,	KΕ,	LS,	MW,	SD,	SL,	SZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,	DE,	DK,
		ES,	FΙ,	FR,	GB,	GR,	IE,	ΙT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,
		CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,	ΤG					
CA	2332	178			A1		2000	0113	1	CA 1	999-	2332	178		1	9990	630
AU	9943	947			Α		2000	0124		AU 1	999-	4394	7		1	9990	630
JP	2000	0800	86		А		2000	0321		JP 1	999-	1864	79		1	9990	630
BR	9911	752			Α		2001	0403		BR 1	999-	1175	2		1	9990	630
ΕP	1092	711			A1		2001	0418		EP 1	999-	9268	53		1	9990	630
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	FI														

TR 200100349	Т2	20010723	TR	2001-200100349		19990630
HU 200102470	A2	20020429	HU	2001-2470		19990630
IN 2000KN00643	A	20050311	IN	2000-KN643		20001218
ZA 2000007635	A	20020102	ZA	2000-7635		20001219
MX 2000PA12925	A	20010521	MX	2000-PA12925		20001220
NO 2000006667	A	20010228	ИО	2000-6667		20001227
LV 12633	В	20010720	LV	2000-177		20001228
US 6545009	B1	20030408	US	2000-720644		20001228
PRIORITY APPLN. INFO.:			JP	1998-186698	Α	19980701
			WO	1999-JP3520	W	19990630

OTHER SOURCE(S): MARPAT 132:93338

GΙ



AB The title compds. I [R1 represents optionally substituted aromatic hydrocarbyl or heteroaryl; R2 represents hydrogen or optionally substituted hydrocarbyl; X represents O, S, etc.; A represents optionally substituted aromatic hydrocarbyl or heteroaryl; and R3 represents OR5, etc.; R5 = H, (un)substituted hydrocarbyl] are prepared I are useful as preventives and remedies for diabetes, etc. Formulations containing I are given. 4-[4-(4-Trifluoromethylphenyl)-2-oxazolyl]benzoic acid at 0.01% in feed decreased blood sugar by 51% in diabetic mice.

IC ICM C07D263-30 ICS C07D277-30; C07D413-04; C07D417-04; A61K031-42; A61K031-425; A61K031-44

CC 28-20 (Heterocyclic Compounds (More Than One Hetero Atom))
Section cross-reference(s): 1, 63

REFERENCE COUNT: 54 THERE ARE 54 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L88 ANSWER 20 OF 20 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1999:736671 ZCAPLUS Full-text

DOCUMENT NUMBER: 131:351319

TITLE: Oxazolylmethoxybenzyl oxyiminoalkanoic acid derivatives with hypoglycemic and hypolipidemic

activity

INVENTOR(S): Momose, Yu; Odaka, Miroyuki; Imoto, Hiroshi;

Kimura, Biroyuki; Sakamoto, Junichi

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Japan

SOURCE: PCT Int. Appl., 148 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND DATE	E APPLICATI	ON NO.	DATE
WO 9958510	A1 1999	01118 WO 1999-J	TP2407	19990510
W: AE, AL, A	M, AU, AZ, BA,	BB, BG, BR, BY,	CA, CN, CU,	CZ, EE, GD,
GE, HR, H	U, ID, IL, IN,	IS, JP, KG, KR,	KZ, LC, LK,	LR, LT, LV,
MD, MG, M	K. MN. MX. NO.	NZ. PL. RO. RU.	SG. SI. SK.	SL. TJ. TM.

					US,														
	RW:	GH,	GM,	KΕ,	LS,	ΜW,	SD,	SL,	SZ,	UC	G, ZV	W, Z	ΑT,	BE,	CH,	СҮ	, DE,	DK,	
		ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC	C, NI	L, I	PT,	SE,	BF,	ВJ	, CF,	CG,	
		CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	ΝE,	SI	J, TI	D, 3	ΤG						
AU	9936297			A					AU 1999-36297						19990510				
	766831				В2														
BR	9910371				Α					BR 1999-10371						19990510			
	1077957				A1												19990510		
EP	1077				В1		2004												
	R:			CH,	DE,	DK,	ES,	FR,	GB,	GF	R, I	Τ,]	LI,	LU,	NL,	SE	, MC,	PT,	
		ΙE,																	
	200003299				Т2											19990510			
	200103714															19990510			
	508066				A					NZ 1999-508066									
	2213738				C2					RU 2000-131183						19990510			
EP	1428				A1		2004				2004						19990		
	R:				DE,	DK,	ES,	FR,	GB,	GF	R, I	T, 1	LI,	LU,	NL,	SE	, MC,	PT,	
			FI,	CY															
	2726				T	20040815				AT 1999-918355						19990510			
	1077				T	20041029				PT 1999-918355						19990510			
	2226				T3					ES 1999-918355 JP 1999-130543						19990510			
	JP 2000034266				A B2	20000202				JP	1999	9-1:	305	43			19990	511	
	JP 3074532						2000				400			0.0					
	JP 2000198772						2000				1999-373202					19990			
	US 6251926						2001			US 1999-423854						19991115			
IN 2000KN00434					A					IN 2000-KN434 MX 2000-PA10576						20001024			
MX 2000PA10576					A									5/6		20001027			
ZA 2000006121					A					ZA 2000-6121 LV 2000-148						20001030			
LV 12606					В					NO 2000-5531						20001101 20001102			
NO 2000005531					A D1		2001			ИО	2000	0-5:	33 I				20001	102	
NO 317426 US 6495581					B1		2004			TTC	2000	Λ 7·	1 1 6 0	20			20001	116	
HK 1034972					B1 A1		20022005										20001		
	US 2003186985						2003			HK 2001-105750 US 2002-331056						20010815 20021227			
	6924		83		A1 B2		2005			0.5	2002	Z-3.	3 T U:	06			20021	221	
	•	DΖ		2005	0002		TD	1998	Q_1	279	21		A	19980	511				
PRIORITY APPLN. INFO.:											1998						19980		
											1999						19990		
											1999						19990		
											1999						19990		
											1999						19991		
											2000						20001		
OTHER SOURCE(S):					MADD	ΔТ	131.	3512°	1 9	00	2000	U — /.	T 40;))	1	. د.	~ O O O I	T T O	
OTHER 9(התעעב.	ΛT	TOT:	$\cup \cup \perp \cup$.	1)													

$$(CH_{2})_{p} = 0 - N = \stackrel{R^{2}}{\stackrel{\cdot}{c}} (CH_{2})_{q} - (\stackrel{R^{4}}{\stackrel{\cdot}{c}})_{m} - \stackrel{\cdot}{\stackrel{\cdot}{c}} = R^{3}$$

$$\stackrel{A}{=} Y - (CH_{2})_{n} - X - R^{1}$$

$$Me$$

$$CH_{2} = 0 - N = \stackrel{\cdot}{\stackrel{\cdot}{c}} - (CH_{2})_{2} - CO_{2}H$$

GI

$$\begin{array}{c} \text{Me} \\ \text{O} \\ \text{Ph} \end{array}$$

$$\text{CH}_2 - \text{O} - \text{N} = \begin{array}{c} \text{Ph} \\ \text{CH}_2 - \text{O} - \text{N} = \begin{array}{c} \text{CH}_2 \end{array}$$

$$\text{CO2H}$$

ΙI

AΒ Title compds. (I) [where R1 = (un)substituted hydrocarbon or heterocyclic group; X = bond, CO, CH(OH), or (alkyl)amino; n = 1-3; Y = 0, S, SO, SO2, or (alkyl)amino; ring A = optionally substituted with 1-3 substituents; p = 1-8; R2 = H or (un)substituted hydrocarbon or heterocyclic group; q = 0-6; m = 0 or 1; R3 = OH, alkoxy, or (un) substituted NH2; R4 and R5 = independently H, hydrocarbon, or may form a ring with R2] were prepared for the prevention or treatment of diabetes mellitus, hyperlipemia, insulin insensitivity, insulin resistance, and impaired glucose tolerance. Thus, reaction of Me (E)-4hydroxyimino-4-phenylbutyrate (preparation given) with 4-(4chloromethylphenoxymethyl)-5-methyl-2-phenyloxazole (preparation given) in DMF followed by deesterification yielded (E)-II (60%). Representative compds. including II were mixed with a powdery diet and fed freely to KKAy mice for 4days. Anal. of blood samples revealed 36% to 54% hypoglycemic action and 35% to 82% hypotriglyceridemic action of the treatment group compared to control animals. Compds. of the invention also exhibited excellent PPARy-RXRlphaheterodimer ligand activity with EC50 values ranging from $0.024~\mu\mathrm{M}$ to $0.79~\mu\mathrm{M}$. TC ICM C07D263-32 ICS A61K031-42; C07C251-54; A61K031-185; C07D413-12; C07D239-42; C07D471-04; C07D413-04; C07D261-08; C07D277-24; C07D215-14; C07D271-06; C07D213-74; C07D471-04; C07D235-00; C07D221-00 CC 28-6 (Heterocyclic Compounds (More Than One Hetero Atom)) Section cross-reference(s): 1 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 4 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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http://www.cas.org/support/stngen/stndoc/properties.html

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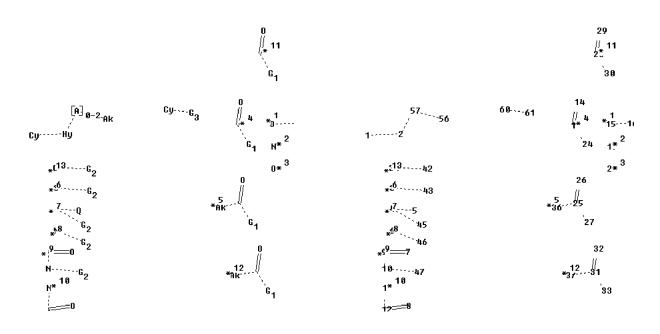
'OBI' IS DEFAULT SEARCH FIELD FOR 'ZCAPLUS' FILE

=> d stat que L72 L12 775523 SEA FILE=REGISTRY ABB=ON PLU=ON N2C3/ES OR NOC3/ES L13 30896 SEA FILE=REGISTRY ABB=ON PLU=ON NSC3/ES L14 805906 SEA FILE=REGISTRY ABB=ON PLU=ON L12 OR L13 L15 464 SEA FILE=REGISTRY ABB=ON PLU=ON NPC3/ES L16 806370 SEA FILE=REGISTRY ABB=ON PLU=ON (L13 OR L14 OR L15)

L19 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation: Uploading L19b.str



chain nodes :

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 20 24 25 26 27 28 29 30 31 32 33 34 36 37 42 43 45 46 47 48 56 57 60 61

ring/chain nodes :

16 17

chain bonds :

 $15-16 \quad 25-26 \quad 25-27 \quad 25-36 \quad 28-29 \quad 28-30 \quad 31-32 \quad 31-33 \quad 31-37 \quad 34-42 \quad 56-57 \quad 60-61 \\ \text{exact/norm bonds:}$

15-16 25-26 25-27 25-36 28-29 28-30 31-32 31-33 31-37 34-42 56-57 60-61

G1:[*1],[*2],[*3]

G2:[*4],[*5]

G3:[*6],[*7],[*8],[*9],[*10],[*11],[*12],[*13]

Connectivity:

2:2 M minimum RC ring/chain 20:1 E exact RC ring/chain

Match level :

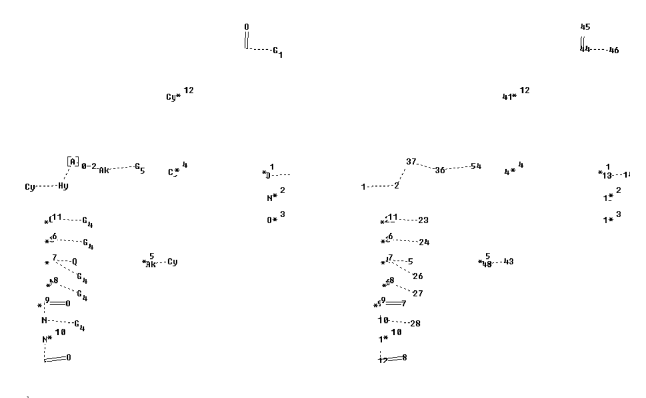
1:Atom 2:Atom 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS 10:CLASS

11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS 20:CLASS

24:CLASS 25:CLASS 26:CLASS 27:CLASS 28:CLASS 29:CLASS 30:CLASS 31:CLASS 32:CLASS 33:CLASS 34:CLASS 36:CLASS 37:CLASS 42:CLASS 43:CLASS 45:CLASS 46:CLASS 47:CLASS 48:CLASS 56:CLASS 57:CLASS 60:Atom 61:CLASS Generic attributes : 60: Saturation : Unsaturated Type of Ring System : Monocyclic Element Count : Node 2: Limited N, N1-2 0,00-1 S,S0-1 P, P0-1 С,С3

L21 71084 SEA FILE=REGISTRY SUB=L16 SSS FUL L19
L23 STR
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

Structure attributes must be viewed using STN Express query preparation: Uploading L23b.str



chain nodes:
1 2 3 4 5 6 7 8 9 10 11 12 13 18 22 23 24 26 27 28 29 36 37

```
41 42 43 44 45 46 48 54
ring/chain nodes :
14 15
chain bonds :
1-2 \quad 2-37 \quad 3-24 \quad 4-5 \quad 4-26 \quad 6-27 \quad 7-9 \quad 8-12 \quad 9-10 \quad 10-28 \quad 11-12 \quad 12-29 \quad 13-14 \quad 22-19 \quad 13-14 \quad 23-19 \quad 13-19 \quad 
23
36-37 36-54 43-48 44-45 44-46
exact/norm bonds :
1-2 \quad 2-37 \quad 3-24 \quad 4-5 \quad 4-26 \quad 6-27 \quad 7-9 \quad 8-12 \quad 9-10 \quad 10-28 \quad 11-12 \quad 12-29 \quad 13-14 \quad 22-19 \quad 13-14 \quad 23-19 \quad 13-14 \quad 23-19 \quad 13-19 \quad 
36-37 36-54 43-48 44-45 44-46
G1:[*1],[*2],[*3]
G4:[*4],[*5]
G5: [*6], [*7], [*8], [*9], [*10], [*11], [*12]
Connectivity:
2:2 M minimum RC ring/chain 18:1 E exact RC ring/chain
Match level:
1:Atom 2:Atom 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS
10:CLASS
11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 18:CLASS 22:CLASS 23:CLASS
24:CLASS 26:CLASS
27:CLASS 28:CLASS 29:CLASS 36:CLASS 37:CLASS 41:Atom 42:Atom 43:Atom
44:CLASS 45:CLASS
46:CLASS 48:CLASS 54:CLASS
Generic attributes :
41:
Saturation
                                                                                                                                                   : Unsaturated
Type of Ring System
                                                                                                                                                       : Monocyclic
                                                                                                                                              : Unsaturated
Saturation
                                                                                                                                       : Monocyclic
Type of Ring System
43:
Saturation
                                                                                                                                                        : Unsaturated
                                                                                                                                        : Monocyclic
Type of Ring System
Element Count :
Node 2: Limited
                     N.N1-2
                        0.00-1
                        S,S0-1
                         P, P0-1
                          C,C3
```

```
L25 31522 SEA FILE=REGISTRY SUB=L21 SSS FUL L23 L29 STR
```

Structure diagram not available for display

Structure attributes must be viewed using STN Express query preparation: Uploading L29b.str

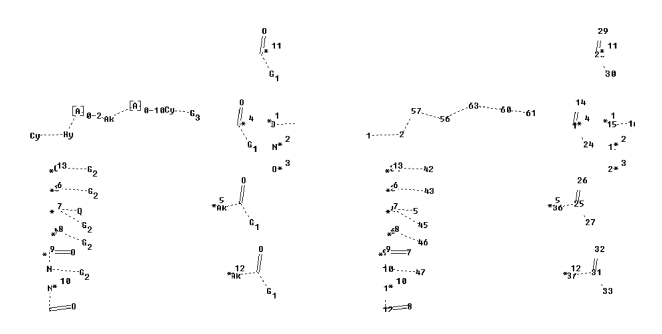
```
42
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            41---- ьз
                                                                                                                                                                                           Cy* <sup>12</sup>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            3{* <sup>12</sup>
                                                                                                                                                                                         CJ* 4
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             3'* <sup>4</sup>
                                                                                                                                                                                                                                                                                                                                                                                                                                     *36-----51
                                                                          cy-----Hy------G
                                                                                                                                                                                                                                                                                                                                                                          1-----83
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             1* 2
                                                                                                                                                                                                                                                                            N* 2
                                  *_0^{14}-Ak<sup>--G</sup>5 *_{c}^{11}-....<sub>G4</sub>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            1* 3
                                                                                                                                                                                                                                                                                                                                    *52°-63--64
                                                                                                                                                                                                                                                                            0*3
                                                                                                                                                                                                                                                                                                                                                                                           *211----23
                                 *5---Ak** G5
                                                                                                                                                                                                                                                                                                                                   *58--65~-66
                                                                                          *:6.....24
                                     *--Q
                                                                                                                                                                                                                                                                                                                                    167--68
5*--55
                                                                                                                                                                                                                                                                                                                                                                                            *1<sup>7</sup><~5
                                                                                           * <sup>7</sup>∴0
                                                                                                                                                                        *<sub>AR</sub>- --Cy
                                                                                                                                                                                                                                                                                                                                                                                                                                                                          *45--40
                                                                                         *<sup>58</sup> - 27
                                  *N---AK--G5
                                                                                                                                                                                                                                                                                                                                    *56--69--70
                                 * <u>18</u>=0
                                                                                                                                                                                                                                                                                                                                    *<sup>5</sup>′-≕61
                                                                                     <u>.9</u>___n
                                                                                                                                                                                                                                                                                                                                                                                       *§<sup>9</sup>—7
                                                                                                                                                                                                                                                                                                                                        57--74--72
                                                                                                                                                                                                                                                                                                                                                                                        10----28
                                      N-Ak G5
                                                                                                                                                                                                                                                                                                                                                                                         1* <sup>10</sup>
                                  *I.19
                                                                                                                                                                                                                                                                                                                                    <sub>*1</sub>19
                                                                                                                                                                                                                                                                                                                                                                                           12===8
                                                                                                                                                                                                                                                                                                                                        66=62
                                          ==0
                                              Ak--Gc
                                                                                                                                                                                                                                                                                                                                               79--74
chain nodes :
1 \quad 2 \quad 3 \quad 4 \quad 5 \quad 6 \quad 7 \quad 8 \quad 9 \quad 10 \quad 11 \quad 12 \quad 13 \quad 18 \quad 22 \quad 23 \quad 24 \quad 26 \quad 27 \quad 28 \quad 29 \quad 36 \quad 38
39 40 41 42 43 45 51 52 53 54 55 56 57 58 59 60 61 62 63 64 65
66 67 68 69
70 71 72 73 74
                                                                                                                                      8.3
ring/chain nodes :
14 15
chain bonds :
1-2 \quad 2-83 \quad 3-24 \quad 4-5 \quad 4-26 \quad 6-27 \quad 7-9 \quad 8-12 \quad 9-10 \quad 10-28 \quad 11-12 \quad 12-29 \quad 13-14 \quad 22-19 \quad 13-14 \quad 23-19 \quad 13-19 \quad 
23
36-51 40-45 41-42 41-43 52-63 53-65 54-55 54-67 56-69 57-59 57-71 58-60
59-61 60-62
60-73 63-64 65-66 67-68 69-70 71-72 73-74
exact/norm bonds :
1-2 \quad 2-83 \quad 3-24 \quad 4-5 \quad 4-26 \quad 6-27 \quad 7-9 \quad 8-12 \quad 9-10 \quad 10-28 \quad 11-12 \quad 12-29 \quad 13-14 \quad 22-19 \quad 13-14 \quad 23-19 \quad 13-14 \quad 23-19 \quad 13-14 \quad 23-19 \quad 13-14 \quad 23-19 \quad 13-19 \quad 
23
36-51 40-45 41-42 41-43 52-63 53-65 54-55 54-67 56-69 57-59 57-71 58-60
59-61 60-62
60-73 63-64 65-66 67-68 69-70 71-72 73-74
G1:[*1],[*2],[*3]
G4:[*4],[*5]
G5:[*6],[*7],[*8],[*9],[*10],[*11],[*12]
G6:[*13],[*14],[*15],[*16],[*17],[*18],[*19]
```

```
Connectivity:
2:2 M minimum RC ring/chain 18:1 E exact RC ring/chain 36:2 E exact RC ring/chain
45:2 E exact RC ring/chain 63:2 E exact RC ring/chain 65:2 E exact RC ring/chain
67:2 E exact
RC ring/chain 69:2 E exact RC ring/chain 71:2 E exact RC ring/chain 73:2 E exact
RC ring/chain
Match level:
1:Atom 2:Atom 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS
10:CLASS
11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 18:CLASS 22:CLASS 23:CLASS
24:CLASS 26:CLASS
27:CLASS 28:CLASS 29:CLASS 36:CLASS 38:Atom 39:Atom 40:Atom 41:CLASS
42:CLASS 43:CLASS
45:CLASS 51:CLASS 52:CLASS 53:CLASS 54:CLASS 55:CLASS 56:CLASS 57:CLASS
58:CLASS 59:CLASS
60:CLASS 61:CLASS 62:CLASS 63:CLASS 64:CLASS 65:CLASS 66:CLASS 67:CLASS
68:CLASS 69:CLASS
70:CLASS 71:CLASS 72:CLASS 73:CLASS 74:CLASS 83:CLASS
Generic attributes :
38:
Saturation
                     : Unsaturated
Type of Ring System
                   : Monocyclic
39:
Saturation
                     : Unsaturated
Type of Ring System
                     : Monocyclic
40:
Saturation
                     : Unsaturated
Type of Ring System
                   : Monocyclic
Element Count:
Node 2: Limited
   N, N1-2
   0.00-1
   S,S0-1
   P, P0-1
   C,C3
```

Structure attributes must be viewed using STN Express query preparation: Uploading L46b.str

L43 16848 SEA FILE=REGISTRY SUB=L25 SSS FUL L29 L46 STR

^{*} STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *



chain nodes :

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 20 24 25 26 27 28 29 30 31 32 33 34 36 37 42 43 45 46 47 48 56 57 60 61 63 ring/chain nodes:

16 17

chain bonds :

15-16 25-26 25-27 25-36 28-29 28-30 31-32 31-33 31-37 34-42 56-57 56-63 60-61 60-63

exact/norm bonds :

G1:[*1],[*2],[*3]

G2:[*4],[*5]

G3:[*6],[*7],[*8],[*9],[*10],[*11],[*12],[*13]

Connectivity:

2:2 M minimum RC ring/chain 20:1 E exact RC ring/chain

Match level:

1:Atom 2:Atom 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS 10:CLASS

11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS 20:CLASS 24:CLASS 25:CLASS

26:CLASS 27:CLASS 28:CLASS 29:CLASS 30:CLASS 31:CLASS 32:CLASS 33:CLASS

```
34:CLASS
36:CLASS 37:CLASS 42:CLASS 43:CLASS 45:CLASS 46:CLASS 47:CLASS 48:CLASS
56:CLASS 57:CLASS
60:Atom 61:CLASS 63:CLASS
Generic attributes :
60.
Saturation : Unsaturated Type of Ring System : Monocyclic
Element Count :
Node 2: Limited
    N.N1-2
    0.00-1
    S, S0-1
    P, P0-1
    C,C3
L48
          8395 SEA FILE=REGISTRY SUB=L43 SSS FUL L46
          3169 SEA FILE=REGISTRY ABB=ON PLU=ON L48 AND NRS<4
L49
          1312 SEA FILE=REGISTRY ABB=ON PLU=ON L49 AND 16.165.12/RID
L51
           785 SEA FILE=REGISTRY ABB=ON PLU=ON 16.167.5/RID AND L49
L57
          2091 SEA FILE=REGISTRY ABB=ON PLU=ON L51 OR L57
L58
L63
              5 SEA FILE=REGISTRY ABB=ON PLU=ON 16.171.9/RID AND L49
          2096 SEA FILE=REGISTRY ABB=ON PLU=ON L58 OR L63
L64
          383 SEA FILE=ZCAPLUS ABB=ON PLU=ON L64
           108 SEA FILE=ZCAPLUS ABB=ON PLU=ON L65 AND J/DT
L66
           275 SEA FILE=ZCAPLUS ABB=ON PLU=ON L65 AND P/DT
L67
            26 SEA FILE=ZCAPLUS ABB=ON PLU=ON L66 AND PY<2003
78 SEA FILE=ZCAPLUS ABB=ON PLU=ON L67 AND PD<20020524
L68
L69
L72
           104 SEA FILE=ZCAPLUS ABB=ON PLU=ON L68 OR L69
=> s L72 not L88
         102 L72 NOT L88
L89
=> d ibib abs hitstr L89 1-102
L89 ANSWER 1 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2003:592703 ZCAPLUS <u>Full-text</u>
DOCUMENT NUMBER:
                        140:13993
TITLE:
                        A new strobin-type fungicide pyraclostrobin
                        Hou, Chunqing; Li, Zhinian; Liu, Changling
AUTHOR(S):
CORPORATE SOURCE:
                       Shenyang Research Inst. of Chemical Industry,
                         Shenyang, 110021, Peop. Rep. China
SOURCE:
                         Nongyao (2002), 41(6), 41-43, 34
                         CODEN: NONGFP; ISSN: 1006-0413
PUBLISHER:
                         Nongyao Bianjibu
DOCUMENT TYPE:
                         Journal; General Review
LANGUAGE:
                         Chinese
     A review on a new strobin-type fungicide pyraclostrobin. It introduces the
     physiochem. properties, toxicity, preparation, mechanism and safety, patent
     and application of pyraclostrobin.
     175013-18-0, Pyraclostrobin
     RL: AGR (Agricultural use); BSU (Biological study, unclassified); BIOL
     (Biological study); USES (Uses)
```

(new strobin-type fungicide)

RN 175013-18-0 ZCAPLUS

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

L89 ANSWER 2 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2003:169613 ZCAPLUS Full-text

DOCUMENT NUMBER: 138:397549

TITLE: Anti-oxidative and anti-senescence effects of the

strobilurin pyraclostrobin in plants: A new strategy

to cope with environmental stress in cereals

AUTHOR(S): Jabs, T.; Pfirrmann, J.; Schafer, S.; Wu, Y. X.; von

Tiedemann, A.

CORPORATE SOURCE: Agricultural Centre, Global Research Biology, BASF AG,

Limburgerhof, 67114, Germany

SOURCE: BCPC Conference--Pests & Diseases (2002),

(Vol. 2), 941-946 CODEN: BCDCAE

PUBLISHER: British Crop Protection Council

DOCUMENT TYPE: Journal LANGUAGE: English

AΒ In addition to its broad spectrum fungicidal activity, the strobilurin pyraclostrobin had pos. effects on the crop yield in the absence of pathogen challenge. This physiol. effect on the plants was especially apparent under conditions of environmental stress. We have observed that pyraclostrobin prevented both symptom development and yield reduction by physiol. leaf spot in barley. Foliar application of pyraclostrobin reduced the production of reactive oxygen intermediates in barley leaf tissues by more than 50% and activated the plant antioxidative system. In addition, pyraclostrobin treatment prevented the release of stress-induced ethylene and premature senescence. Since the physiol. leaf spot disease and other environmental stresses are caused by changes in the genetic and metabolic regulation of reactive oxygen intermediates resulting in membrane-leakage, cell death or premature senescence, we postulate that the anti-oxidative and anti-senescence effects of pyraclostrobin are responsible for its ability to improve stress tolerance in plants.

IT *175013-18-0*, Pyraclostrobin

RL: AGR (Agricultural use); BSU (Biological study, unclassified); BIOL (Biological study); USES (Uses)

(anti-oxidative and anti-senescence effects of pyraclostrobin in barley)

RN 175013-18-0 ZCAPLUS

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 11 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 3 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2003:169607 ZCAPLUS Full-text

DOCUMENT NUMBER: 138:349947

TITLE: Shift in sensitivity of Alternaria solani (potato

early blight) to strobilurin fungicides

AUTHOR(S): Pasche, J. S.; Wharam, C. M.; Gudmestad, N. C. Department of Plant Pathology, North Dakota State CORPORATE SOURCE:

University, Fargo, ND, 58105, USA

SOURCE: BCPC Conference--Pests & Diseases (2002),

> (Vol. 2), 841-846 CODEN: BCDCAE

PUBLISHER: British Crop Protection Council

DOCUMENT TYPE: Journal LANGUAGE: English

Forty-seven Alternaria solani isolates collected from 1998 and 2001 from AB various potato growing regions in the United States were assayed in vitro for sensitivity to azoxystrobin. Twenty-one A. solani isolates collected in 1998, prior to the introduction of azoxystrobin, had a mean baseline EC50 value of 0.0279µq/mL. Isolates of A. solani collected in 2001, recovered from fields displaying a lack of disease control by azoxystrobin, had a mean EC50 of $0.3480 \mu g/mL$. Mean EC50 values for baseline isolates to pyraclostrobin and trifloxystrobin were $0.0022\mu g/mL$ and $0.0060 \mu g/mL$ resp. In 2001, sensitivities to pyraclostrobin and trifloxystrobin shifted to mean EC50 values of $0.0208\mu q/mL$, and $0.0140\mu q/mL$ resp. In vivo assessments of sensitivity to azoxystrobin and pyraclostrobin were conducted on six isolates selected from the in vitro cross-resistance evaluations. Results from the in vivo assays were correlated to those obtained in the in vitro assays. Field studies need to be conducted to determine if the shift in sensitivity to pyraclostrobin and trifloxystrobin will result in a similar loss of disease control under com. potato growing conditions as observed with azoxystrobin.

TΤ 175013-18-0, Pyraclostrobin

> RL: BSU (Biological study, unclassified); BIOL (Biological study) (sensitivity of Alternaria solani (potato early blight) to strobilurin fungicides)

RN 175013-18-0 ZCAPLUS

CMCarbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

REFERENCE COUNT: 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 4 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:824680 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 138:319985

TITLE: Pyraclostrobin; pesticide tolerance

CORPORATE SOURCE: Environmental Protection Agency, Office of Pesticide

Programs, Environmental Protection Agency, Washington,

DC, 20460, USA

SOURCE: Federal Register (2002), 67(188),

60886-60902, 27 Sep 2002

CODEN: FEREAC; ISSN: 0097-6326 Superintendent of Documents

DOCUMENT TYPE: Journal LANGUAGE: English

PUBLISHER:

Tolerances are established for combined residues of pyraclostrobin (carbamic acid, [2-[[[1-(4-chlorophenyl)-1H-pyrazol-3- yl]oxy]methyl]phenyl]methoxy-, Me ester) and its desmethoxy metabolite Me 2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl carbamate, expressed as parent compound, in or on almond, hulls and various other fruits and vegetables and agricultural products, and combined residues of pyraclostrobin and its metabolites convertible to 1-(4-chlorophenyl)-1H-pyrazol-3-ol, expressed as parent compound, in or on cattle, fat and various other animal products. BASF Corporation requested these tolerances under the Federal Food, Drug, and Cosmetic Act (FFDCA), as amended by the Food Quality Protection Act (FQPA) of 1996.

IT 175013-18-0, Pyraclostrobin 512165-96-7

RL: ADV (Adverse effect, including toxicity); POL (Pollutant); BIOL (Biological study); OCCU (Occurrence)

(tolerance for pyraclostrobin of food and feed)

RN 175013-18-0 ZCAPLUS

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-

yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

RN 512165-96-7 ZCAPLUS

CN Carbamic acid, [2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

L89 ANSWER 5 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:717832 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 137:334223

TITLE: A strobilurin fungicide enhances the resistance of

tobacco against tobacco mosaic virus and Pseudomonas

syringae pv tabaci

AUTHOR(S): Herms, Stefan; Seehaus, Kai; Koehle, Harald; Conrath,

Uwe

CORPORATE SOURCE: Department of Biology, University of Kaiserslautern,

Kaiserslautern, D-67653, Germany

SOURCE: Plant Physiology (2002), 130(1), 120-127

CODEN: PLPHAY; ISSN: 0032-0889

PUBLISHER: American Society of Plant Biologists

DOCUMENT TYPE: Journal LANGUAGE: English

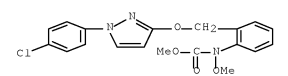
A strobilurin fungicide, F 500 (Pyraclostrobin), enhances the resistance of AΒ tobacco (Nicotiana tabacum cv Xanthi nc) against infection by either tobacco mosaic virus (TMV) or the wildfire pathogen Pseudomonas syringae pv tabaci. F 500 was also active at enhancing TMV resistance in NahG transgenic tobacco plants unable to accumulate significant amts. of the endogenous inducer of enhanced disease resistance, salicylic acid (SA). Apparently, F 500 enhances TMV resistance in tobacco either by acting downstream of SA in the SA signaling mechanism or by functioning independently of SA. The latter assumption is the more likely because in infiltrated leaves, F 500 did not cause the accumulation of SA-inducible pathogenesis-related (PR)-1 proteins that often are used as conventional mol. markers for SA-induced disease resistance. However, accumulation of PR-1 proteins and the associated activation of the PR-1 genes were elicited upon TMV infection of tobacco leaves and both these responses were induced more rapidly in F 500-pretreated plants than in the water-pretreated controls. Thus, F 500, in addition to exerting direct antifungal activity, may also protect plants by priming them for potentiated activation of subsequently pathogen-induced cellular defense responses.

IT 175013-18-0, Pyraclostrobin

RL: BSU (Biological study, unclassified); BIOL (Biological study) (F 500 strobilurin fungicide enhancement of resistance of tobacco against tobacco mosaic virus and Pseudomonas syringae tabaci)

RN 175013-18-0 ZCAPLUS

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)



REFERENCE COUNT: 28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 6 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:602334 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 137:347825

TITLE: Efficiency of fungicides to control anthracnose and

angular leaf spot in common beans

AUTHOR(S): Rava, Carlos A.

CORPORATE SOURCE: Embrapa Arroz e Feijao, Santo Antonio de Goias, Brazil

SOURCE: Summa Phytopathologica (2002), 28(1), 65-69

CODEN: SUPHDV; ISSN: 0100-5405

PUBLISHER: Grupo Paulista de Fitopatologia

DOCUMENT TYPE: Journal LANGUAGE: Portuguese

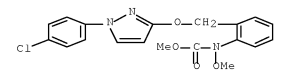
The effect of spray applications of two active ingredients, alone and in AΒ mixts.: epoxyconazole to control angular leafspot and pyraclostrobin, to control both anthracnose and angular leaf spot of common beans was studied. The treatments tested for control of anthracnose were. carbendazim + epoxyconazole (250 + 12.5 g ha-1); thiophanate Me + epoxyconazole (300 + 12.5 g ha-1); pyraclostrobin (50, 75, 100 g ha-1); pyraclostrobin + epoxyconazole $(26.6 + 10 \ 33.3 + 12.5 \ g \ ha-1)$; tebuconazole $(200 \ g \ ha-1)$; and the check. For the angular leaf spot control trial, besides the above treatments were also included: epoxyconazole (12.5 g ha-1); azoxystrobin (60 g ha-1); tebuconazole (200 g ha-1); and thiophanate Me + chlorothalonil (350+875 g ha-1). Pyraclostrobin alone or in mixture with epoxyconazole, significantly reduced anthracnose severity, in all tested doses. All fungicides and doses tested to control anthracnose increased grain yield significantly, reaching as much as 97% increase in comparison with the check. Epoxyconazole alone or in mixts., showed high efficiency for control angular leaf spot. The effect of pyraclostrobin in all three doses tested and its mixture with epoxyconazole did not differ from epoxyconazole alone and in mixture with carbendazim and thiophanate. These treatments showed significantly higher control efficiency of angular leaf spot than azoxystrobin, tebuconazole and thiophanate Me + chorothalonil.

IT 175013-18-0, Pyraclostrobin

RL: BSU (Biological study, unclassified); BIOL (Biological study) (fungicides for control anthracnose and angular leaf spot in common beans)

RN 175013-18-0 ZCAPLUS

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)



REFERENCE COUNT: 28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 7 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:353222 ZCAPLUS Full-text

DOCUMENT NUMBER: 136:351654

TITLE: Polymeric pest control sheet containing pesticides

INVENTOR(S):
Barazani, Avner

PATENT ASSIGNEE(S): Makhteshim Chemical Works Ltd., Israel

SOURCE: PCT Int. Appl., 21 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:	TENT NO.	K	KIND DATE		APPLICAT	ION NO.	DATE				
WO		Ž	A2 20020	510 W	VO 2001-	 IL1014		20011101 <			
	W: AE, AG CO, CR GM, HR LS, LT PL, PT UG, US RW: GH, GM	AL, AI, CU, CU, HU, III, LU, LV, RO, RU, UZ, VI, KE, LS	M, AT, AU, Z, DE, DK, D, IL, IN, V, MA, MD, U, SD, SE, N, YU, ZA,	AZ, BA, DM, DZ, IS, JP, MG, MK, SG, SI, ZW SD, SL,	EC, EE, KE, KG, MN, MW, SK, SL, SZ, TZ,	ES, FI, KP, KR, MX, MZ, TJ, TM,	GB, GI KZ, LO NO, NZ TR, TI	D, GE, GH, C, LK, LR, Z, OM, PH, I, TZ, UA, E, CH, CY,			
	·	•	I, CM, GA,	GN. GO.	GW. MI.	MR. NE.	SN. TI				
EG CA AU EP	139388 22884 2427485 200214232 1330160	1 1 1	A 20050 A 20031 A1 20020 A 20020 A2 20030	925 I 030 E 510 C 515 <i>P</i>	IL 2000-1 IG 2001-1 IA 2001-1 AU 2002-1	139388 157 2427485 14232		20001101 20011030 20011101 < 20011101 < 20011101			
		CH, DI			GR, IT,	LI, LU,	NL, SE	SE, MC, PT,			
BR JP AT RU AU ES IN NO BG ZA	200302985 2001015377 2004513896 336892 2292136 2002214232 2271087 2003KN00516 2003001858 107766 2003003283 2003PA03889		A 20040 I 20040 I 20060 C2 20070 B2 20070 A 20041 A 20030 A 20040 A 20030	128	EU 2003-: BR 2001-: BR 2001-: AT 2001-: AU 2003-:	15377 538753 982691 112457 214232 1982691 KN516 1858 107766 3283 PA3889		20011101 20011101 20011101 20011101 20011101 20030424 20030425 20030429 20030429 20030430			
	2004025413 Y APPLN. INF). :	A1 20040					20030501 A 20001101			
					WO 2001-IL1014			W 20011101			

AB A sheet for pest control is made of polymeric material and comprises at least two layers; a top layer and a bottom layer, wherein the bottom layer contains a herbicide and one or more pesticides selected from among fungicides and insecticides, and the top layer optionally contains an insecticide and/or fungicide. Other aspects of the invention include a polymeric composition used in the preparation of the sheets and a method for pest control in agriculture, horticulture and gardens.

IT *175013-18-0*, Pyraclostrobin

RL: AGR (Agricultural use); BSU (Biological study, unclassified); BIOL (Biological study); USES (Uses)

(polymeric pest control sheet containing)

RN 175013-18-0 ZCAPLUS

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

L89 ANSWER 8 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:312686 ZCAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER: 137:30451

TITLE: Sensitivity to azoxystrobin among isolates of Uncinula

necator: baseline distribution and relationship to

myclobutanil sensitivity

AUTHOR(S): Wong, Francis P.; Wilcox, Wayne F.

CORPORATE SOURCE: Department of Plant Pathology, New York Agricultural

Experiment Station, Cornell University, Geneva, NY,

14456, USA

SOURCE: Plant Disease (2002), 86(4), 394-404

CODEN: PLDIDE; ISSN: 0191-2917

PUBLISHER: American Phytopathological Society

DOCUMENT TYPE: Journal LANGUAGE: English

Two hundred fifty-six single-conidial chain isolates of Uncinula necator were assayed for their sensitivity to azoxystrobin and myclobutanil. These isolates were collected from two sites in New York in 1999: an organic vineyard where no synthetic fungicides have been used (baseline population) and a com. vineyard having a history of compromised powdery mildew control with myclobutanil (demethylation inhibitor [DMI]-resistant population). Mean coeffs. of variance for a leaf disk assay used to test fungicide sensitivities were 31% for azoxystrobin and 41% for myclobutanil. Baseline ED50 values ranged from 0.0037 to $0.028 \, \mu \text{g/mL}$ (mean $0.0097 \, \mu \text{g/mL}$) for azoxystrobin and from 0.0049 to $0.69~\mu\text{g/mL}$ (mean $0.075~\mu\text{g/mL}$) for myclobutanil. A shift in the mean ED50 value for azoxystrobin to $0.018~\mu g/mL$ was observed in the DMIresistant population; with the strongest shift observed for isolates collected from vines treated exclusively with myclobutanil (0.024 $\mu g/mL$). For the 256 tested isolates, there was a moderate, but statistically significant, correlation between azoxystrobin and myclobutanil sensitivities (R2=0.36, P<0.001). Tests with three other strobilurin fungicides (kresoxim-Me, pyraclostrobin, and trifloxystrobin) indicate clear differences in the intrinsic activity of these compds. against U. necator, and the applicability of the methods developed with azoxystrobin for assays with pyraclostrobin and trifloxystrobin. Isolates from the high and low ends of the azoxystrobin sensitivity distribution (15x difference in mean ED50 values) were equally controlled in planta by protectant or postinfection treatment with azoxystrobin at 250 μg a.i./mL, but postinfection application at lower rates (2.5 and 25 μq a.i./mL) resulted in a 41 and 44% decrease, resp., in the control of the low-sensitivity isolates vs. high-sensitivity isolates. The results of this study document the baseline sensitivity distribution of U. necator to azoxystrobin, provide evidence of partial cross-sensitivity between azoxystrobin and myclobutanil, and illustrate the potential selection for individuals with reduced sensitivity (quant. range) to azoxystrobin by postinfection application and reduced rates of this fungicide.

IT *175013-18-0*, Pyraclostrobin

RL: AGR (Agricultural use); BSU (Biological study, unclassified); BIOL (Biological study); USES (Uses)

(strobilurin fungicide sensitivity among Uncinula necator isolates)

RN 175013-18-0 ZCAPLUS

N Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

REFERENCE COUNT: 42 THERE ARE 42 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 9 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:293365 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 136:320810

TITLE: Synergistic insecticidal, fungicidal and acaricidal

mixtures

INVENTOR(S): Fischer, Reiner; Wachendorff-Neumann, Ulrike

PATENT ASSIGNEE(S): Bayer Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 79 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Fatent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.					KIND DATE			APPLICATION NO.						DATE 			
WO	2002	0301	- -		A1		2002	0418	,	WO 2	001-	 EP11	 126		2	00109	926 <
	W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PH,	PL,
		PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG,
		US,	UZ,	VN,	YU,	ZA,	ZW										
	RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,
		DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,	TR,	BF,
		ВJ,	CF,	CG,	CI,	CI, CM, GA, GN,				GW,	ML,	MR,	NE,	SN,	TD,	ΤG	
DE	10049	9804			A1		2002	0418		DE 2	000-	1004	9804		2	0001	009 <
AU	2002	1396	7		А		2002	0422	AU 2002-13967						2	0109	926 <
ΕP	1326	495			A1		2003	0716	EP 2001-982360				20010926				
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR						
BR	2001	01449	91		Α		2003	1014		BR 2	001-	1449	1		2	0109	926
JΡ	2004	51079	93		Τ		2004	0408	8 JP 2002-533652					2	0109	926	
									4 IN 2001-MU931					20010926			
									27 US 2003-398265						403		
MΧ	20031	PA03	029	A 2003062					24 MX 2003-PA3029				20030407				
ORITY APPLN. INFO.:								DE 2	000-	1004	9804	2	A 2	20001009			
									,	WO 2	001 - 1	EP11	126	Ī	W 2	00109	926

OTHER SOURCE(S): MARPAT 136:320810

AB The title mixts. comprise known cyclic ketoenole (Markush given) and any of 55 known insecticides, fungicides or acaricides, such as fluquinconazole, tebuconazole, bitertanol, triadimenol, triadimefon, difenoconazole, flusilazole, prochloraz, penconazole, etc.

IT 175013-18-00, BAS 500F, mixts. with cyclic ketoenol derivs.

RL: AGR (Agricultural use); BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)

(synergistic pesticidal mixts.)

RN 175013-18-0 ZCAPLUS

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 10 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:240497 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 136:279449

TITLE: Preparation of 4-acylaminopyrazole derivatives as

agrochemicals

INVENTOR(S): Kajino, Hisaki; Morimoto, Munetsugu; Furuta, Satoru;

Tanaka, Hisako; Tanaka, Harukazu; Ohnishi, Tohru

PATENT ASSIGNEE(S): Sankyo Company, Ltd., Japan

SOURCE: PCT Int. Appl., 985 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA'	PATENT NO.						KIND DATE			APPLICATION NO.						DATE 				
WO	2002	0239	86		A2		2002	0328	,	WO 2	001-	JP71	66		2	0010	821	<		
	W:	ΑE,	AG,	AL,	ΑM,	ΑT,	ΑU,	AZ,	BA,	BB,	ВG,	BR,	BY,	BZ,	CA,	CH,	CN,			
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,			
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	ΚE,	KG,	KR,	KΖ,	LC,	LK,	LR,	LS,	LT,			
		LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MΖ,	NO,	NΖ,	PH,	PL,	PT,	RO,			
		RU,	SD,	SE,	SG,	SI,	SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,			
		VN,	YU,	ZA,	ZW,	ΑM,	ΑZ,	BY,	KG,	KΖ,	MD,	RU,	ТJ,	TM						
	RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,			
		DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,			
		ΒJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	ΤG				
AU	2001	0800	99		A5		2002	0402		AU 2	001-	8009	9		2	0010	821	<		
EP	1329	160			A2		2003	0723		EP 2	001-	9583	83		2	0010	821			
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙT,	LI,	LU,	NL,	SE,	MC,	PT,			
		IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR									
JP	JP 2002138082						2002	0514	1	JP 2	001-	2523	48		2	0010	823	<		
PRIORIT	RIORITY APPLN. INFO.:								1	JP 2	000-	2548	09		A 2	0000	825			
												WO 2001-JP7166					W 20010821			
OTHER SO	THER SOURCE(S):					MARPAT 136:27944				449										

AB The title compds. I [R1 is hydrogen, optionally substituted C1-16 alkyl, or the like; R2 and R3 are each independently hydrogen, halogeno, optionally substituted C1-6 alkyl, or the like; R4 is hydrogen, C1-6 alkyl, or cyano; Z is oxygen or sulfur; Ar is optionally substituted C6-14 aryl or an optionally substituted 5- or 6-membered unsatd. heterocyclic group; and B is hydrogen, halogeno, optionally substituted C1-16 alkyl, or the like] are prepared Me N-(3-cyanobenzyl)-N-(1-isobutyl-3-methyl-1H- pyrazole)carbamate at 10 ppm gave ≥ 50% control of Phytophthora infestans.

IT 405545-51-9P 405545-53-1P 405545-54-2P 405545-55-3P 405545-56-4P 405545-57-5P 405545-58-6P 405545-59-7P 405545-60-0P 405545-68-8P 405546-72-7P

RL: AGR (Agricultural use); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of 4-acylaminopyrazole derivs. as agrochems.) $405545-51-9\ \text{ZCAPLUS}$

CN Benzeneacetic acid, 2-[[acetyl(3,5-dimethyl-1-phenyl-1H-pyrazol-4-yl)amino]methyl]- α -(methoxymethylene)-, methyl ester (9CI) (CA INDEX NAME)

RN

RN 405545-53-1 ZCAPLUS
CN Benzoic acid, 3-[[acetyl(3,5-dimethyl-1-phenyl-1H-pyrazol-4-yl)amino]methyl]- (9CI) (CA INDEX NAME)

RN 405545-54-2 ZCAPLUS

CN Benzoic acid, 3-[[acetyl(3,5-dimethyl-1-phenyl-1H-pyrazol-4-yl)amino]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 405545-55-3 ZCAPLUS

CN Benzoic acid, 3-[[acetyl(3,5-dimethyl-1-phenyl-1H-pyrazol-4-yl)amino]methyl]-, ethyl ester (9CI) (CA INDEX NAME)

CN Benzoic acid, 3-[[acetyl(3,5-dimethyl-1-phenyl-1H-pyrazol-4-yl)amino]methyl]-, 1-methylethyl ester (9CI) (CA INDEX NAME)

RN 405545-57-5 ZCAPLUS

CN Benzamide, 3-[[acetyl(3,5-dimethyl-1-phenyl-1H-pyrazol-4-yl)amino]methyl]- (9CI) (CA INDEX NAME)

RN 405545-58-6 ZCAPLUS

CN Benzamide, 3-[[acetyl(3,5-dimethyl-1-phenyl-1H-pyrazol-4-yl)amino]methyl]-N-methyl- (9CI) (CA INDEX NAME)

RN 405545-59-7 ZCAPLUS

CN Benzamide, 3-[[acetyl(3,5-dimethyl-1-phenyl-1H-pyrazol-4-yl)amino]methyl]-N-propyl-(9CI) (CA INDEX NAME)

RN 405545-60-0 ZCAPLUS

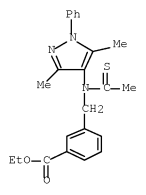
CN Benzamide, 3-[[acetyl(3,5-dimethyl-1-phenyl-1H-pyrazol-4-yl)amino]methyl]-N-(cyanomethyl)- (9CI) (CA INDEX NAME)

RN 405545-68-8 ZCAPLUS

CN Benzamide, 3-[[acetyl(3,5-dimethyl-1-phenyl-1H-pyrazol-4-yl)amino]methyl]-N-(1-oxopropyl)- (9CI) (CA INDEX NAME)

RN 405546-72-7 ZCAPLUS

CN Benzoic acid, 3-[[(3,5-dimethyl-1-phenyl-1H-pyrazol-4-yl)(1-thioxoethyl)amino]methyl]-, ethyl ester (9CI) (CA INDEX NAME)



L89 ANSWER 11 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:234509 ZCAPLUS Full-text

DOCUMENT NUMBER: 137:93732

TITLE: Synthesis of new salicylamide derivatives with

evaluation of their antiinflammatory, analgesic and

antipyretic activities

AUTHOR(S): Fahmy, H. H.; Soliman, G. A.

CORPORATE SOURCE: Therapeutical Chemistry Department, National Research

Centre, Cairo, Egypt

SOURCE: Archives of Pharmacal Research (2001),

24(3), 180-189

CODEN: APHRDQ; ISSN: 0253-6269

PUBLISHER: Pharmaceutical Society of Korea

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 137:93732

AB A new series of pyridazine, pyrazoles, pyrazolidine-3,5-dione, semicarbazide, thiosemicarbazides, hydantoin, thiohydantoins, 1,2,4-triazoles, S-triazolo[3,4-b]-1,3,4-thiadiazoles incorporated indirectly into salicylamide moiety at position 2 were synthesized. Also the synthesis of novel series of 3-salicylamido-2-hydroxypropyl amine derivs. were prepared Several of these compds. were screened for antiinflammatory, analgesic, antipyretic and ulcerogenic activities.

IT 442129-55-7P

RN 442129-55-7 ZCAPLUS

CN Benzamide, 2-[2-(5-amino-3-phenyl-1H-pyrazol-1-yl)-2-oxoethoxy]- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 12 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:220302 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 136:243290

TITLE: Synergistic fungicidal compositions INVENTOR(S): Nuninger, Cosima; Zeller, Martin PATENT ASSIGNEE(S): Syngenta Participations A.-G., Switz.

SOURCE: PCT Int. Appl., 33 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT	ATENT NO.				KIND DATE			APPLICATION NO.						DATE 			
WO 2002	0219	18		A1		2002	0321		WO 2	001-	EP10	446		2	0010	910 <-	
W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	AZ,	BA,	BB,	ВG,	BR,	BY,	BZ,	CA,	CH,	CN,	
	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	
	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,	
	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PH,	PL,	
	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	
	US,	UZ,	VN,	YU,	ZA,	ZW											
RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,	
	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,	TR,	BF,	
	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG		
CA 2421	226			A1		2002	0321		CA 2	001-	2421.	226		2	0010	910 <-	
AU 2002	1222	7		А		2002	0326		AU 2	002-	1222	7		2	0010	910 <-	
EP 1317	178			A1		2003	0611		EP 2	001-	9803	66		2	0010	910	
EP 1317	178			В1		2004	0512										
R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,	

	IE,	SI, LT,	LV,	FI, RO, MK,	CY, Al	L, TR		
BR :	200101381	.5	А	20030708	BR	2001-13815		20010910
HU :	200301024	:	A2	20031028	HU	2003-1024		20010910
AT :	266316		T	20040515	AT	2001-980366		20010910
JP :	200451862	:3	T	20040624	JP	2002-526185		20010910
TW :	220381		В	20040821	TW	2001-90122367		20010910
ES :	2217194		Т3	20041101	ES	2001-1980366		20010910
RU :	2270564		C2	20060227	RU	2003-109610		20010910
ZA :	200300156	9	А	20040420	ZA	2003-1569		20030226
IN :	2003CN003	48	А	20050408	IN	2003-CN348		20030307
MX :	2003PA021	17	A	20030619	MΧ	2003-PA2117		20030311
US :	200318995	8	A1	20031009	US	2003-380486		20030312
PRIORITY	APPLN. I	NFO.:			GB	2000-22338	A	20000912
					WO	2001-EP10446	W	20010910

OTHER SOURCE(S): MARPAT 136:243290

GΙ

- AB The title compns. comprise a N-sulfonylvalinamide I (R1 = H, C1-4 alkyl, C3-6 cycloalkyl or halophenyl; R2 = C1-4 alkyl) in association with acibenzolar-S-Me, azoxystrobin, chlorothalonil, cymoxanil, dimethomorph, fluazinam, fludioxonil, imazalil, S-imazalil, mancozeb, metalaxyl, metalaxyl-M, picoxystrobin, pyraclostrobin (BAS 500F), trifloxystrobin, etc.
- IT 175013-18-0D, Pyraclostrobin, mixts. with N-sulfonylvalinamide derivs.
 - RL: AGR (Agricultural use); BIOL (Biological study); USES (Uses) (synergistic fungicidal compns.)
- RN 175013-18-0 ZCAPLUS
- CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-vl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 13 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:106491 ZCAPLUS Full-text DOCUMENT NUMBER: 136:351622

TITLE: Evaluation of fungicides in control of spot-type net

blotch on barley

AUTHOR(S): Jayasena, K. W.; Loughman, R.; Majewski, J.

CORPORATE SOURCE: Agriculture Western Australia, Albany, 6330, Australia

SOURCE: Crop Protection (2002), 21(1), 63-69

CODEN: CRPTD6; ISSN: 0261-2194

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

Ten fungicides (pyraclostrobin, tebuconazole, flutriafol, epoxiconazole, propiconazole, triadimefon, azoxystrobin, trifloxystrobin, difenoconazole and a mixture of propiconazole with iprodione) were evaluated as single applications for control of spot-type net blotch of barley caused by Drechslera teres maculata at three locations during 1999 and 2000. Under moderate disease severity, yield losses ranged from 17-19% depending on location and under high disease severity, yield losses reached 32%. Pyraclostrobin, propiconazole and a mixture of propiconazole with iprodione were the most effective in controlling disease, improving yield and grain quality. These fungicides show most promise as com. treatments when yield and quality are taken into account. Azoxystrobin, trifloxystrobin, difenoconazole and epoxiconazole also provided disease control.

IT *175013-18-0*, (Pyraclostrobin

RL: BSU (Biological study, unclassified); BIOL (Biological study) (fungicides in control of spot-type net blotch on barley)

RN 175013-18-0 ZCAPLUS

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ \end{array}$$

REFERENCE COUNT: 19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 14 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:851100 ZCAPLUS Full-text

DOCUMENT NUMBER: 135:371520

TITLE: Preparation of novel phenyl propargyl ethers as

agrochemical fungicides

INVENTOR(S): Lamberth, Clemens; Zeller, Martin; Kunz, Walter;

Cederbaum, Fredrik

PATENT ASSIGNEE(S): Syngenta Participations A.-G., Switz.

SOURCE: PCT Int. Appl., 84 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAI	PATENT NO.				KIND DATE				APPL	ICAT	ION I		DATE				
WO	WO 2001087822 W: AE, AG, AI				A1	_	2001	 1122		 WO 2	001-	EP55		20010515 <			
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         TW 228117
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                                            A1 20011122 CA 2001-2406088
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        AU 2001260301
BR 2001010810
                                           B2 20041104
                                            A 20030211 BR 2001-10810
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                                            A 20030211 BR 2001-10810 20010515
A1 20030212 EP 2001-933967 20010515
         EP 1282595
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         EP 1282595
                R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
                       IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
         HU 200301965 A2 20030929
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      AT 271031
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      AT 2001-933967

      PT 1282595
      T
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      ES 2223848
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      ES 2001-1933967

      RU 2259353
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      RU 2002-133216

      EG 22695
      A
      20030630
      EG 2001-511

      IN 2002CN01841
      A
      20050211
      IN 2002-CN1841

      MX 2002PA11198
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      20030310
      MX 2002-PA11198

      ZA 200209266
      A
      20031020
      ZA 2002-9266

      US 6683211
      B1
      20040127
      US 2002-276476

      HR 2002000908
      B1
      20060731
      HR 2002-908

      HK 1054368
      A1
      20050603
      HK 2003-104881

      RTTY APPLN. INFO::
      GB 2000-11944

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                                                                               HK 2002-908 20021115

HK 2003-104881 20030708

GB 2000-11944 A 20000517

WO 2001-E05530 W 20010515

WO 2001-EP5530 W 20010515
PRIORITY APPLN. INFO.:
OTHER SOURCE(S): MARPAT 135:371520
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$$R^{1}-C \equiv C \xrightarrow{R^{2}} O \xrightarrow{R^{4}O} \xrightarrow{R^{5}} \xrightarrow{R^{7}} \underbrace{H} \xrightarrow{O} \overset{Z}{\xrightarrow{P}} R^{1}O$$

GΙ

RN

175013-18-0 ZCAPLUS

The title compds. [I; R1 = H, alkyl, cycloalkyl, (un)substituted aryl; R2, R3 = H, alkyl; R4 = alkyl, alkenyl, alkynyl; R5- R8 = H, alkyl; R9 = H, (un)substituted alkyl, alkenyl or alkynyl; R10 = (un)substituted (hetero)aryl; Z = halo, (un)substituted aryloxy, alkoxy, etc.] which possess useful plant protecting properties and may advantageously be employed in agricultural practice for controlling or preventing the infestation of plants by phytopathogenic microorganisms, especially fungi (biol. data given), were prepared E.g., a multi-step synthesis of I [R1-R3 = H; R4 = Me; R5-R8 = H; R9 = H; R10 = 4-ClC6H4; Z = OMe] was given.

IT 175013-18-0, Pyraclostrobin RL: AGR (Agricultural use); BIOL (Biological study); USES (Uses) (preparation of novel Ph propargyl ethers as agrochem. fungicides)

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 15 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:816378 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 135:340474

TITLE: Method for inducing antiviral resistance in plants

INVENTOR(S): Koehle, Harald; Conrath, Uwe; Seehaus, Kai

PATENT ASSIGNEE(S): BASF Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 26 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

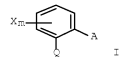
FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

									APPLICATION NO.										
	2001															0010	 430 <-	_	
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		HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,	LS,		
		LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,		
		RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,		
		VN,	YU,	ZA,	ZW														
	RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,		
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CA	2409	649			A1		2001	1108		CA 2	001-	2409	649		2	0010	430 <-	_	
EP	1278	415			A1		2003	0129		EP 2	001-	9472	50		2	0010	430		
EP	1278	415			В1		2003	1001											
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BR	2001	0104	55		A		2003	0311		BR 2	001-	1045	5		2	0010	430		
HU	2003	0063	1		A2		2003	0728		HU 2	003-	631			2	0010	430		
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JP	2003	5318														0010	430		
PT	1278	415					2004									0010			
NZ	5223	41			Α		2004	0430								0010			
	2210				Т3		2004	0701		ES 2	001-	1947	250		2	0010	430		
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	2003						2003					2578							
	2002						2003					PA10			2	0021	025		
	2002						2003					9751				0021			
	2004				A1		2004	0923				8169				0040			
PRIORIT	Y APP	LN.	INFO	.:								1002							
											EP48								
									US 2	002-	2578	74		A1 2	0021	017			
OTHER S	R SOURCE(S):				MARI	PAT	PAT 135:340			Į									

OTHER SOURCE(S): MARPAT 135:340474

GΙ



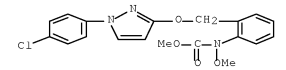
The invention relates to a method for inducing antiviral resistance in plants, which is characterized in that the plants, the soil or seeds are treated with a compound, which is absorbed by the plants or seeds. The compds. are I [X = halo, C1-4 alkyl or trifluoromethyl; m = 0 or 1; Q = C(:CHCH3)COOCH3, C(:CHOCH3)COOCH3, C(:CHOCH3)CONHCH3 or N(OCH3)COOCH3, C(:CHOCH3)CONHCH3, C(:NOCH3)COOCH3, C(:NOCH3)CONHCH3 or N(OCH3)COOCH3; A = OB, CH2OB, OCH2B, CH:CHB, C:CB, CH2ON:CR1B or CH2ON:CR1CR2:NOR3; B = (un)substituted Ph, naphthyl, 5-member or 6-member heteroaryl or 5-member or 6-member heterocyclyl, containing one to three N atoms and/or one O or S atom or one or two O and/or S atoms; R1 = H, cyano, alkyl, haloalkyl, cycloalkyl or alkoxy; R2 = (un)substituted Ph, phenylcarbonyl, phenylsulfonyl, 5-member or 6-member heteroarylsulfonyl, etc.; R3 = H or (un)substituted alkyl, alkenyl and alkynyl]. Prefered I are pyraclostrobin, picoxystrobin, trifloxystrobin and azoxystrobin.

IT 175013-18-0, Pyraclostrobin

RL: AGR (Agricultural use); BIOL (Biological study); USES (Uses) (inducing antiviral resistance in plants by)

RN 175013-18-0 ZCAPLUS

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)



REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 16 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:781491 ZCAPLUS Full-text

DOCUMENT NUMBER: 136:69768

TITLE: Design, Synthesis, and Biological Evaluation of a

Library of 1-(2-Thiazolyl)-5-(trifluoromethyl)pyrazole-

4-carboxamides

AUTHOR(S): Donohue, Bridget A.; Michelotti, Enrique L.; Reader,

John C.; Reader, Valerie; Stirling, Matthew; Tice,

Colin M.

CORPORATE SOURCE: Rohm and Haas Company, Spring House, PA, 19477-0904,

USA

SOURCE: Journal of Combinatorial Chemistry (2002),

4(1), 23-32

CODEN: JCCHFF; ISSN: 1520-4766

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

AΒ A library of 422 1-(2-thiazoly1)-5-(trifluoromethy1)pyrazole-4- carboxamides was prepared in five steps using solution-phase chemical The first step in the synthesis was the reaction of Et 2-ethoxymethylene-3-oxo-4,4,4trifluorobutanoate with thiosemicarbazide, which is reported in the literature to afford a 1:1 mixture of Et 1-thiocarbamoyl-5- (trifluoromethyl)pyrazole-4carboxylate and Et 1-thiocarbamoyl-3- (trifluoromethyl)pyrazole-4-carboxylate. The product is, however, a single compound, Et 5-hydroxy-1-thiocarbamoyl-5-(trifluoromethyl)-4,5- dihydro-1H-pyrazole-4-carboxylate. This common intermediate was diversified by reaction with 17 α -bromo ketones affording, in two steps, 17 1-(2-thiazoly1)-5-(trifluoromethy1)pyrazole-4-carboxylic acids. Scavenger resins were used to facilitate formation and purification of up to 27 amides from each of these acids in the last step. In addition, the Curtius reaction was applied to 12 of the acids followed by quenching with alcs. to afford a 108-member carbamate library. Certain compds. in the two libraries were toxic to C. elegans.

IT 385412-59-9P 385412-60-2P

RL: CPN (Combinatorial preparation); CMBI (Combinatorial study); PREP (Preparation)

(preparation and nematocidal activity of a library of 1-(2-thiazolyl)-5-(trifluoromethyl) pyrazole-4-carboxamides and -carbamates)

RN 385412-59-9 ZCAPLUS

CN Benzoic acid, 4-[[[[1-(4-ethyl-2-thiazolyl)-5-(trifluoromethyl)-1H-pyrazol-4-yl]carbonyl]amino]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 385412-60-2 ZCAPLUS

CN Benzoic acid, 4-[[[[1-[4-(1,1-dimethylethyl)-2-thiazolyl]-5-(trifluoromethyl)-1H-pyrazol-4-yl]carbonyl]amino]methyl]-, methyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT: 41 THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 17 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:780351 ZCAPLUS Full-text

DOCUMENT NUMBER: 135:299954

TITLE: Fungicidal compositions comprising

methoxyiminoacetamide derivatives.
INVENTOR(S): Wachendorff-Neumann, Ulrike; Seitz, Thomas; Gayer,

Machendorfi-Neumann, Ulrike; Seitz, Inomas; Gayer,
Herbert; Heinemann, Ulrich; Krueger, Bernd-Wieland;

Kraemer, Wolfgang; Assmann, Lutz

PATENT ASSIGNEE(S): Bayer A.-G., Germany SOURCE: Ger. Offen., 40 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Fatent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

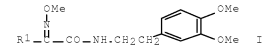
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PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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EP 1276375 EP 1276375	A2 B1	20030122 20050720	W, ML, MR, NE, SN, T EP 2001-933807 BB, GR, IT, LI, LU, N	20010409
BR 2001010116 JP 2003531154 HU 200302686 AT 299648	T A2 T	20030211 20031021 20031128 20050815	EY, AL, TR BR 2001-10116 JP 2001-577751 HU 2003-2686 AT 2001-933807 PT 2001-933807	20010409 20010409 20010409

ES 2	2243496	T3	20051201	ES	2001-1933807		20010409
RU 2	2265331	C2	20051210	RU	2002-131167		20010409
IN 2	2001MU00339	A	20050304	IN	2001-MU339		20010412
ZA 2	2002007474	A	20030918	ZA	2002-7474		20020918
US 2	2003158151	A1	20030821	US	2002-257740		20021016
US 6	6787567	B2	20040907				
MX 2	2002PA10331	A	20030523	MX	2002-PA10331		20021018
US 2	2004266850	A1	20041230	US	2004-840907		20040507
PRIORITY	APPLN. INFO.:			DE	2000-10019758	Α	20000420
				WO	2001-EP4042	W	20010409
				US	2002-257740	АЗ	20021016

OTHER SOURCE(S): MARPAT 135:299954

GΙ



- AB Fungicidal compns. comprise methoxyiminoacetamide derivs. I (R1 = fluorine-, chlorine-, bromine-, Me-, Et-, Pr- iso-Pr-, Bu-, iso-Bu-, tert-Bu-, methoxy-, ethoxy- or phenoxy-substituted or unsubstituted Ph, 2-naphthyl, 1,2,3,4-tetrahydronaphthyl, indanyl, 2-benzofuranyl, 2-benzothienyl, 2-thienyl or 2-furanyl) and any of known 58 fungicides.
- IT 175013-18-0D, mixts. with methoxyiminoacetamide derivs.
 RL: AGR (Agricultural use); BIOL (Biological study); USES (Uses)
 (fungicidal compns.)
- RN 175013-18-0 ZCAPLUS
- CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

L89 ANSWER 18 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:747733 ZCAPLUS Full-text

DOCUMENT NUMBER: 135:303727

TITLE: Synthesis of lunularic acid derivatives as

chemopreventive agents

INVENTOR(S): Gerhaeuser, Clarissa; Eicher, Theophil; Pick, Rigobert

PATENT ASSIGNEE(S): Deutsches Krebsforschungszentrum Stiftung Des

Oeffentlichen Rechts, Germany

SOURCE: PCT Int. Appl., 64 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Fatent
LANGUAGE: German
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

P	PATENT NO.					KIND DATE		APPLICATION NO.					DATE					
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	LV, MA, MD SE SG SI			MD,	MG,	MK,	MN,	MW,	MX,	MΖ,	NO,	NΖ,	PL,	PT,	RO,	RU,	SD,	
	SE, SG, SI			SI,	SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,	YU,	
	ZA, ZW, AM			ΑM,	ΑZ,	BY,	KG,	KΖ,	MD,	RU,	ТJ,	TM						
	RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,	
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		ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG			
D:							2001	1011		DE 2	000-	1001	5525		2	0000	330 <-	
PRIORI'	PRIORITY APPLN. INFO.:									DE 2000-10015525					A 20000330			
OTHER	OTHER SOURCE(S):				MARPAT 135:3037													
GI	` '																	

$$R^3$$
 CO_2R^2 I

Lunularic acid derivs. [I-IV; X = (un)substituted mono or polycyclic (hetero)aryl; R1, R2 = alkyl, alkenyl, mono or polycyclic aryl; R3 = F, Cl, Br, I, amino. alkylamino, aminoalkyl, OH, carboxyl, alkoxycarbonyl, carbamoyl, aryl, acyloxy, etc.] are prepared which are suitable as chemopreventive agents. Thus, lunularic acid derivative II [R1 = R3 = H, R2 = Me, X = Ph (V)] was prepared via Wittig reaction between (3-acetoxy-2-methoxycarbonyl)benzyl-triphenyl-phosphonium bromide and benzaldehyde. V was tested for chemopreventive properties (IC50 = 0.087 μM vs. CyplA1 in Hepalc1c7 mouse hepatoma cells; 40% inhibition of DMBA-induced preneoplastic lesions in mice thymus gland culture; IC50 = 7.2 μM for inhibition of quinone oxidoreductase induction).

IT 365542-56-9P 365542-57-0P 365542-58-1P 365542-59-2P 365542-60-5P 365542-61-6P 365542-74-1P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological

RN 365542-56-9 ZCAPLUS

CN Benzoic acid, 2-[(1E)-2-(5-chloro-3-methyl-1-phenyl-1H-pyrazol-4-yl)ethenyl]-6-methoxy-, methyl ester (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 365542-57-0 ZCAPLUS

CN Benzoic acid, 2-[2-(5-chloro-3-methyl-1-phenyl-1H-pyrazol-4-yl)ethyl]-6-methoxy-, methyl ester, monohydrochloride (9CI) (CA INDEX NAME)

● HCl

RN 365542-58-1 ZCAPLUS

CN Benzoic acid, 2-[(1E)-2-(5-chloro-3-methyl-1-phenyl-1H-pyrazol-4-yl)ethenyl]-6-methoxy-, ethyl ester (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 365542-59-2 ZCAPLUS

CN Benzoic acid, 2-[2-(5-chloro-3-methyl-1-phenyl-1H-pyrazol-4-yl)ethenyl]-6-methoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 365542-60-5 ZCAPLUS

CN Benzoic acid, 2-[(1Z)-2-(5-chloro-3-methyl-1-phenyl-1H-pyrazol-4-yl)ethenyl]-6-methoxy-, methyl ester, monohydrochloride (9CI) (CA INDEX NAME)

Double bond geometry as shown.

● HCl

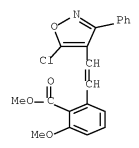
RN 365542-61-6 ZCAPLUS

CN Benzoic acid, 2-[2-(5-chloro-3-methyl-1-phenyl-1H-pyrazol-4-yl)ethenyl]-6-methoxy-, methyl ester, monohydrochloride (9CI) (CA INDEX NAME)

● HCl

RN 365542-74-1 ZCAPLUS

CN Benzoic acid, 2-[2-(5-chloro-3-phenyl-4-isoxazolyl)ethenyl]-6-methoxy-, methyl ester (9CI) (CA INDEX NAME)



REFERENCE COUNT: 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 19 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:472679 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 135:61328

TITLE: Preparation of 2-[2-(1-phenyl-1H-pyrazol-3-

 $\verb|yl|) oxymethylphenyl]-2-fluoromethoxyiminoacetates and$

methylacetamides as agrochemical fungicides and

insecticides

INVENTOR(S): Heinemann, Ulrich; Gayer, Herbert; Gerdes, Peter;

Krueger, Bernd-Wieland; Maurer, Fritz; Vaupel, Martin;
Mauler-Machnik, Astrid; Wachendorff-Neumann, Ulrike;
Haenszler, Gerd; Kuck, Karl-Heinz; Loesel, Peter;

Erdelen, Christoph

PATENT ASSIGNEE(S): Bayer Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 61 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Fatent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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PATENT NO.
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                                        WO 2000-EP12481
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            HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
            LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
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PRIORITY APPLN. INFO.:
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                                          DE 2000-10034129
                                                            A 20000713
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                                         WO 2000-EP12481
OTHER SOURCE(S):
                      MARPAT 135:61328
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GΙ

$$\begin{array}{c} R1 \\ R2 \\ R4 \\ R4 \\ KMe \end{array}$$

AB Title compds. [I; X = O, NH; R = (substituted) alkyl, cycloalkyl, aryl; R1-R4 = H, halo, cyano, NO2, (substituted) alkyl, alkoxy, alkylthio, alkylsulfinyl, alkylsulfonyl], were prepared Thus, Me 2-[2-[1-(4- chlorophenyl)-1H-pyrazol-3-yl]oxymethylphenyl]-2-fluoromethoxyiminoacetate (preparation given) in MeOH was treated with MeNH2 followed by stirring for 18 h to give 60.5% 2-[2-[1-(4- chlorophenyl)-1H-pyrazol-3-yl]oxymethylphenyl]- 2-fluoromethoxyimino-N-methylacetamide. Tested I at 250 g/ha gave ≥98% control of Erysiphe graminis f.sp. hordei on barley.

Ι

345905-74-0P 345905-75-1P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of phenylpyrazolyloxymethylphenylfluoromethoxyiminoacetates

and

methylacetamides as agrochem. fungicides and insecticides)

RN 345905-38-6 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl] oxy] methyl]- $\alpha-[(\text{fluoromethoxy}) \text{imino}]-, \text{methyl ester (9CI)} (CA INDEX NAME)}$

RN 345905-39-7 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)imino]-N-methyl- (9CI) (CA INDEX NAME)

RN 345905-40-0 ZCAPLUS

CN Benzeneacetamide, α -[(fluoromethoxy)imino]-2-[[[1-(4-fluorophenyl)-1H-pyrazol-3-yl]oxy]methyl]-N-methyl- (9CI) (CA INDEX NAME)

RN 345905-41-1 ZCAPLUS

CN Benzeneacetic acid, α -[(fluoromethoxy)imino]-2-[[[1-[4-(1-methylethyl)phenyl]-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 345905-42-2 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(3-bromophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)imino]-N-methyl- (9CI) (CA INDEX NAME)

RN 345905-43-3 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(4-bromophenyl)-1H-pyrazol-3-yl]oxy]methyl]- $\alpha-[(fluoromethoxy)imino]-N-methyl- (9CI) (CA INDEX NAME)$

RN 345905-44-4 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(3,5-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- $\alpha-[(fluoromethoxy)imino]-N-methyl- (9CI) (CA INDEX NAME)$

RN 345905-45-5 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(2-\text{chlorophenyl})-1H-\text{pyrazol}-3-\text{yl}] \text{oxy}] \text{methyl}]-\alpha-[(fluoromethoxy)imino]-, methyl ester (9CI) (CA INDEX NAME)$

RN 345905-46-6 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(4-cyanophenyl)-1H-pyrazol-3-yl]oxy]methyl]- $\alpha-[(\text{fluoromethoxy})\text{imino}]-, \text{methyl ester (9CI)}$ (CA INDEX NAME)

RN 345905-47-7 ZCAPLUS

CN Benzeneacetic acid, α -[(fluoromethoxy)imino]-2-[[[1-(4-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 345905-48-8 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(2,4-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)imino]-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \operatorname{FCH}_2 - \operatorname{O} - \operatorname{N} & \operatorname{O} \\ & & & \\ \operatorname{C} - \operatorname{C} - \operatorname{OMe} \\ & & & \\ \operatorname{CH}_2 - \operatorname{O} & & \\ \end{array}$$

RN 345905-49-9 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(2-bromophenyl)-1H-pyrazol-3-yl]oxy]methyl]- $\alpha-[(fluoromethoxy)imino]-, methyl ester (9CI) (CA INDEX NAME)$

RN 345905-50-2 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(4-bromophenyl)-1H-pyrazol-3-yl]oxy]methyl]- $\alpha-[(fluoromethoxy)imino]-, methyl ester (9CI) (CA INDEX NAME)$

RN 345905-51-3 ZCAPLUS

CN Benzeneacetic acid, α -[(fluoromethoxy)imino]-2-[[(1-phenyl-1H-pyrazol-3-yl)oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 345905-52-4 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(3,5-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)imino]-, methyl ester (9CI) (CA INDEX NAME)

RN 345905-53-5 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(3-bromophenyl)-1H-pyrazol-3-yl]oxy]methyl]- $\alpha-[(fluoromethoxy)imino]-, methyl ester (9CI) (CA INDEX NAME)$

RN 345905-54-6 ZCAPLUS

CN Benzeneacetic acid, α -[(fluoromethoxy)imino]-2-[[[1-(4-fluorophenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 345905-55-7 ZCAPLUS

CN Benzeneacetamide, α -[(fluoromethoxy)imino]-N-methyl-2-[[(1-phenyl-1H-pyrazol-3-yl)oxy]methyl]- (9CI) (CA INDEX NAME)

RN 345905-56-8 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(2,4-dichlorophenyl)-1H-pyrazol-3-yl]]] $\alpha-[(fluoromethoxy)]$ mino]-N-methyl- (9CI) (CA INDEX NAME)

RN 345905-57-9 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-[4-(1,1-dimethylethyl)phenyl]-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)imino]-, methyl ester (9CI) (CA INDEX NAME)

RN 345905-58-0 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(2-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)imino]-N-methyl- (9CI) (CA INDEX NAME)

RN 345905-59-1 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-[4-(1,1-dimethylethyl)phenyl]-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)imino]-N-methyl- (9CI) (CA INDEX NAME)

RN 345905-60-4 ZCAPLUS

CN Benzeneacetic acid, α -[(fluoromethoxy)imino]-2-[[[1-(2-fluorophenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 345905-61-5 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(2,4-difluorophenyl)-1H-pyrazol-3-yl]oxy]methyl]-<math>\alpha-[(fluoromethoxy)imino]-$, methyl ester (9CI) (CA

INDEX NAME)

RN 345905-62-6 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(2,3-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)imino]-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \operatorname{FCH}_2-\operatorname{O-N} & \operatorname{O} \\ & \operatorname{C-C-OMe} \\ & \operatorname{CH}_2-\operatorname{O-Me} \\ & \operatorname{C1} \end{array}$$

RN 345905-63-7 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(3,4-dimethylphenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)imino]-, methyl ester (9CI) (CA INDEX NAME)

RN 345905-64-8 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(2,5-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)imino]-, methyl ester (9CI) (CA INDEX NAME)

RN 345905-65-9 ZCAPLUS

CN Benzeneacetic acid, α -[(fluoromethoxy)imino]-2-[[[1-(3-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 345905-66-0 ZCAPLUS

CN Benzeneacetamide, α -[(fluoromethoxy)imino]-N-methyl-2-[[[1-(3-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]- (9CI) (CA INDEX NAME)

RN 345905-67-1 ZCAPLUS

CN Benzeneacetamide, α -[(fluoromethoxy)imino]-2-[[[1-(2-fluorophenyl)-1H-pyrazol-3-yl]oxy]methyl]-N-methyl- (9CI) (CA INDEX NAME)

RN 345905-68-2 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(2,4-difluorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)imino]-N-methyl- (9CI) (CA INDEX NAME)

RN 345905-69-3 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(2,3-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)imino]-N-methyl- (9CI) (CA INDEX NAME)

FCH₂
$$\longrightarrow$$
 O N O NHMe CH₂ \longrightarrow CH₂ \longrightarrow O NHMe

RN 345905-70-6 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(2,5-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)imino]-N-methyl- (9CI) (CA INDEX NAME)

FCH₂—O—N O
$$C_1$$
 CH₂—O—NHMe

RN 345905-71-7 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(3-\text{chlorophenyl})-1H-\text{pyrazol}-3-\text{yl}] \text{oxy}] \text{methyl}] - \alpha-[(fluoromethoxy)imino]-, methyl ester (9CI) (CA INDEX NAME)$

RN 345905-72-8 ZCAPLUS

CN Benzeneacetamide, α -[(fluoromethoxy)imino]-N-methyl-2-[[[1-[4-(1-methylethyl)phenyl]-1H-pyrazol-3-yl]oxy]methyl]- (9CI) (CA INDEX NAME)

RN 345905-73-9 ZCAPLUS

CN Benzeneacetamide, α -[(fluoromethoxy)imino]-N-methyl-2-[[[1-(4-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]- (9CI) (CA INDEX NAME)

RN 345905-74-0 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(4-cyanophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)imino]-N-methyl- (9CI) (CA INDEX NAME)

RN 345905-75-1 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(3-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)imino]-N-methyl- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 20 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:452988 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 135:46183

TITLE: Preparation of dihalopropenyloxybenzene derivatives

and pesticides containing the same as the active

ingredient

INVENTOR(S): Katsurada, Manabu; Kawata, Shinji; Kyomura, Nobuo;

Shiga, Yasushi; Fukuchi, Toshiki; Yamada, Risa

PATENT ASSIGNEE(S): Mitsubishi Chemical Corporation, Japan

SOURCE: PCT Int. Appl., 135 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Fatent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

GΙ

PATENT NO.	K	ND D	ATE	А	PPLI	CAT	ON 1	4O.		DZ	ATE	
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OTHER SOURCE(S):	MA	RPAT 1	35:46183	3								

$$A = (CR^{1}R^{2})p - W = (CR^{3}R^{4})q - Q = (CR^{5}R^{6})r$$

$$X^{1} = CZ^{2}$$

$$X^{2} = (CR^{1}R^{2})p - W = (CR^{3}R^{4})q - Q = (CR^{5}R^{6})r$$

AΒ Dihalopropenyloxybenzene derivs. such as (dichloropropenyloxyphenyl)isoxaz ole, (dichloropropenyloxy) benzene, and (dichloropropenyloxyphenyl) oxadiazo le derivs. represented general formula [I; A = H, (un) substituted alkyl, alkenyl, alkynyl, aryl, or heterocyclyl; W = single bond, O, S, SO, SO2, NR7, N:(R7), C(R7):NO, ON:C(R7), C(R7):NN:C(R8), CO, CO2, O2C, N(R7)CO, CON(R7); wherein R7, R8 = H, alkyl; Q = SO, SO2, N:C(R9), C(R9):NO, ON:C(R9), C(R9):N:C(R10), CO, CO2, O2C, N(R9)CO, CON(R9), (un)substituted aryl or heterocyclyl; R9, R10 = H, alkyl; R1 - R6 = H, alkyl, alkenyl, alkynyl, alkoxy, alkenyloxy, alkynyloxy, alkoxyalkyl; or R1 and R2, R3 and R4, or R5 and R6 together form a alkylidene or alkylidenedioxy; p, q, and r are integers and $p+q+r\leq 9$, provided that when Q represents SO, SO2, C(R9):NO, CO2, or CONR9, r is ≥1; when Q represents Ph, r is 0; when Q represents ON:C(R9) or O2C and W represents O or S, q is ≥ 1 ; X1, X2 = H, halo, alkyl, haloalkyl; Y = halo, alkyl, haloalkyl; n = 0-2; Z = halo] are prepared These compds. have a very excellent effect of controlling pests in the field of agriculture, horticulture, foods, clothing, housing, livestock, pets, etc. (in particular, injurious insects and mites in the fields of agriculture and horticulture) and are highly safe to mammals and fishes. Thus, chlorination of 2,6-dichloro-4-(3,3-dichloro-2propenyloxy) benzaldoxime (preparation given) by N-chlorosuccinimide in THF at room temperature for 2 h followed by ammonolysis with NH3 in MeOH at room temperature for 2 h gave 2,6-dichloro-4-(3,3-dichloro-2propenyloxy) benzamidoxime which was treated with NaH at room temperature for 20 min and at 60° for 25 min and cyclocondensed with Et glycolate to give 3-

[2,6-dichloro-4-(3,3-dichloro-2-propenyloxy)phenyl]-5-hydroxymethyl- 1,2,4-oxadiazole. Etherification of the latter alc. with 3-trifluoromethylphenol using PPh3 and di-Et azodicarboxylate in THF at room temperature for 15.5 h gave $3-[2,6-\text{dichloro-4-((3,3-\text{dichloro-2-propenyl)oxy)phenyl]-5-((3-trifluoromethylphenoxy)methyl)-1,2,4-oxadiazole which at 500 ppm controlled 100% larvae of Plutella xylostella Plutella xylostella konaga, Spodoptera litura, and Adoxophyes sp.$

IT 345199-56-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of dihalopropenyloxybenzene derivs. and pesticides containing same

as active ingredient)

RN 345199-56-6 ZCAPLUS

CN Benzoic acid, 4-[2-[3-[2,6-dichloro-4-[(3,3-dichloro-2-propenyl)oxy]phenyl]-5-isoxazolyl]ethoxy]-, methyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT: 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 21 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:449811 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 135:46178

TITLE: Preparation of methyl 2-[2-(1-phenyl-1H-pyrazol-3-

yl)oxymethylphenyl]-3-fluoromethoxy-2-acrylates as agrochemical fungicides, insecticides, and acaricides.

INVENTOR(S): Heinemann, Ulrich; Gayer, Herbert; Gerdes, Peter;

Krueger, Bernd-Wieland; Maurer, Fritz; Vaupel, Martin; Mauler-Machnik, Astrid; Wachendorff-Neumann, Ulrike; Haenssler, Gerd; Kuck, Karl-Heinz; Erdelen, Christoph;

Loesel, Peter

PATENT ASSIGNEE(S): Bayer A.-G., Germany SOURCE: Ger. Offen., 18 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Fatent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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            SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
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                         A1
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                         В1
PRIORITY APPLN. INFO.:
                                           DE 1999-19961330 A 19991220 WO 2000-EP12322 W 20001207
OTHER SOURCE(S):
                       MARPAT 135:46178
GΙ
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AΒ Title compds. [I; R1 = alkyl, cycloalkyl, aryl; R2-R5 = H, halo, cyano, NO2, (halo-substituted) alkyl, alkoxy, alkylthio, alkylsulfinyl, alkylsulfonyl] were prepared Thus, 2-(2-bromoethylphenyl)-3-(fluoromethoxy)- 2-acrylic acid Me ester was stirred for 18 h at room temperature with 1-(4-chlorophenyl)-1,2dihydro-3H-pyrazol-3-one and NaH in DMF to give 48% Me 2-[2-[1-(4chlorophenyl)-1H-pyrazol-3-yl]oxymethylphenyl]-3- fluoromethoxy-2-acrylate. Tested I at 250 g/ha gave 98% control of Puccinia recondita on wheat. 344569-93-3P 344569-94-4P 344569-95-5P 344569-96-6P 344569-97-7P 344569-98-8P 344569-99-9P 344570-00-9P 344570-01-0P 344570-02-1P 344570-03-2P 344570-04-3P 344570-05-4P 344570-06-5P RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of phenylpyrazolyloxymethylphenylfluoromethoxyacrylic acid Me esters as agrochem. fungicides, insecticides, and acaricides) 344569-93-3 ZCAPLUS RN CN Benzeneacetic acid, 2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]-

 α -[(fluoromethoxy)methylene]-, methyl ester (9CI) (CA INDEX NAME)

RN 344569-94-4 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-[4-(1,1-dimethylethyl)phenyl]-1H-pyrazol-3-yl]oxy]methyl]-<math>\alpha-[(fluoromethoxy)methylene]-$, methyl ester (9CI) (CA INDEX NAME)

$$t-Bu$$

$$V$$

$$FCH_2-O-CH=C$$

$$MeO-C$$

RN 344569-95-5 ZCAPLUS

CN Benzeneacetic acid, α -[(fluoromethoxy)methylene]-2-[[[1-(4-fluorophenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 344569-96-6 ZCAPLUS

CN Benzeneacetic acid, α -[(fluoromethoxy)methylene]-2-[[[1-[4-(1-methylethyl)phenyl]-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 344569-97-7 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(3-bromophenyl)-1H-pyrazol-3-yl]oxy]methyl]- $\alpha-[(fluoromethoxy)methylene]-, methyl ester (9CI) (CA INDEX NAME)$

RN 344569-98-8 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(4-bromophenyl)-1H-pyrazol-3-yl]oxy]methyl]- $\alpha-[(fluoromethoxy)methylene]-, methyl ester (9CI) (CA INDEX NAME)$

RN 344569-99-9 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(3,5-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)methylene]-, methyl ester (9CI) (CA INDEX NAME)

RN 344570-00-9 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(2-\text{chlorophenyl})-1H-\text{pyrazol}-3-\text{yl}] \text{oxy}] \text{methyl}] - \alpha-[(fluoromethoxy) \text{methylene}]-, methyl ester (9CI) (CA INDEX NAME)$

RN 344570-01-0 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(4-cyanophenyl)-1H-pyrazol-3-yl]oxy]methyl]- $\alpha-[(\text{fluoromethoxy})\text{methylene}]-, \text{methyl ester (9CI)}$ (CA INDEX NAME)

RN 344570-02-1 ZCAPLUS

CN Benzeneacetic acid, α -[(fluoromethoxy)methylene]-2-[[[1-(4-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{Me} \\ \text{Me} \\ \end{array}$$

RN 344570-03-2 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(2,4-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)methylene]-, methyl ester (9CI) (CA INDEX NAME)

RN 344570-04-3 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(2-bromophenyl)-1H-pyrazol-3-yl]oxy]methyl]- $\alpha-[(fluoromethoxy)methylene]-, methyl ester (9CI) (CA INDEX NAME)$

RN 344570-05-4 ZCAPLUS

CN Benzeneacetic acid, α -[(fluoromethoxy)methylene]-2-[[(1-phenyl-1H-pyrazol-3-yl)oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 344570-06-5 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-[2,6-dichloro-4-(trifluoromethyl)phenyl]-1H-pyrazol-3-yl]oxy]methyl]- α -[(fluoromethoxy)methylene]-, methyl ester (9CI) (CA INDEX NAME)

L89 ANSWER 22 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:338507 ZCAPLUS Full-text

DOCUMENT NUMBER: 134:340502

TITLE: Preparation of benzoylcyclohexanediones and

benzoylpyrazoles as herbicides and plant growth

regulators.

INVENTOR(S):
Seitz, Thomas; Willms, Lothar; Auler, Thomas;

Bieringer, Hermann; Thuerwaechter, Felix

PATENT ASSIGNEE(S): Aventis CropScience GmbH, Germany

SOURCE: PCT Int. Appl., 113 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Fatent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA'	TENT	NO.			KIN:	D	DATE			APPL	ICAT	ION 1	. OV		D.	ATE	
WO	2001	 0326	 36		A1		2001	0510		WO 2	000-	EP10	460		2	0001	024 <
	W:	ΑE,	AG,	AL,	AM,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CN,	CR,	CU,
		CZ,	DM,	DZ,	EE,	GD,	GE,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KG,	ΚP,	KR,
		KΖ,	LC,	LK,	LR,	LT,	LV,	MA,	MD,	MG,	MK,	MN,	MX,	NO,	NΖ,	PL,	RO,
		RU,	SG,	SI,	SK,	ΤJ,	TM,	TR,	TT,	UA,	UZ,	VN,	YU,	ZA,	ΑM,	ΑZ,	BY,
		KG,	KΖ,	MD,	RU,	ΤJ,	$_{ m MT}$										
	RW:	GH,	GM,	ΚE,	LS,	MW,	MΖ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,
		DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,
		CF,	CG,	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,	ΤG			
CA	2397	361			A1		2001	0510		CA 2	000-	2397.	361		2	0001	024 <
BR	2000	0153	38		Α		2002	0723		BR 2	000-	1533	8		2	0001	024
EP	1235	816			A1		2002	0904		EP 2	000-	9744	43		2	0001	024
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙT,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL							
JP	2003	5130	81		Τ		2003	0408		JP 2	001-	5347	87		2	0001	024
US	6448	201			В1		2002	0910		US 2	000-	7050	01		2	0001	102
PRIORIT	Y APP	LN.	INFO	.:						DE 1	999-	1995.	3136		A 1	9991	104
										WO 2	000-	EP10	460		W 2	0001	024
OTHER SO	OURCE	(S):			MAR:	PAT	134:	3405	02								

Ι

$$\mathbb{Q} \xrightarrow{\text{NHetR}^3(\mathbb{R}^4)} \mathbb{Q}$$

$$Q^{1} = (P^{5})$$

$$(R^{6})$$

$$Q^{2} = N$$

$$\mathbb{R}^{7}$$

$$\mathbb{R}^{8}$$

$$\mathbb{R}^{8}$$

AΒ Title compds. [I; Q = Q1, Q2; X = OR3a, OCOR3a, OCONHR3a, OSO2R3a, alkyl, alkenyl, alkynyl, Ph, etc.; R1, R2 = H, SH, NO2, halo, cyano, alkyl, alkoxyalkyl, haloalkyl, alkenyl, alkynyl, etc.; R3 = H, OH, halo, SH, amino, cyano, NO2, CHO, alkoxycarbonyl, alkylcarbonyl, etc.; R3a = H, (substituted) alkyl, alkenyl, alkynyl, Ph, phenylalkyl; R4 = [C(R11)2]mAr[C(R11)2]mR12; A = O, S; R5 = OR16, alkylthio, haloalkylthio, alkenylthio, haloalkenylthio, alkynylthio, haloalkynylthio, alkylsulfinyl, haloalkylsulfinyl, etc.; R6 = H, tetrahydropyranyl, tetrahydrothiopyranyl, (substituted) alkyl, cycloalkyl, alkoxy, alkylcarbonyl, alkoxyalkyl, etc.; R7 = H, alkyl, haloalkyl; R8 = alkyl, haloalkyl, (substituted) Ph; R9 = H, alkyl, haloalkyl, alkylcarbonyl, alkoxycarbonyl, haloalkylcarbonyl, alkoxycarbonyl, alkylsulfonyl, haloalkylsulfonyl, (substituted) PhCO, PhCOCH2, PhOCO2, PhSO2, etc.; R11 = H, alkyl, halo; R12 = (substituted) cycloalkyl, cycloalkenyl, aryl, heterocyclyl, heteroaryl, etc.; Y = O, S, NH, CHR6, C(R6)2, alkylimino; Z = bond, O, S, SO, SO2, NH, alkylimino, CHR7, C(R7)2; m, n = 0-2; p = 1, 2; q = 0-4; r = 0, 1], were prepared Thus, 2-chloro-3-(3-phenylisoxazol-5-yl)methoxy-4methylsulfonylbenzoic acid (preparation given), cyclohexane-1,3-dione, N'-(3dimethylaminopropyl)-N- ethylcarbodiimide hydrochloride, and dimethylaminopyridine were stirred in CH2Cl2 to give 60% enol ether, which was stirred with acetone cyanohydrin, Et3N, and KCN in MeCN to give 55% 2-[2chloro-3-(3-phenylisoxazol-5- yl)methoxy-4-methylsulfonylbenzoyl]cyclohexan-1,3-dione. Several I at ≤ 1 kg/ha postemergent gave $\geq 80\%$ control of Sinapis alba and Stellaria media.

IT 338461-87-3P 338461-88-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of benzoylcyclohexanediones and benzoylpyrazoles as herbicides and plant growth regulators)

RN 338461-87-3 ZCAPLUS

CN Benzoic acid, 2-chloro-4-(methylsulfonyl)-3-[(3-phenyl-5-isoxazolyl)methoxy]-, methyl ester (9CI) (CA INDEX NAME)

RN 338461-88-4 ZCAPLUS

CN Benzoic acid, 2-chloro-4-(methylsulfonyl)-3-[(3-phenyl-5-isoxazolyl)methoxy]- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 11 THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 23 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:338479 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 134:353175

TITLE: Preparation of amides and ureas as activators of

soluble quanylate cyclase

INVENTOR(S): Selwood, David; Glen, Robert; Reynolds, Karen;

Wishart, Grant

PATENT ASSIGNEE(S): University College London, UK

SOURCE: PCT Int. Appl., 101 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA	TENT	NO.			KIN	D	DATE				ICAT				D.	ATE	
WO	2001	0326	04		A1		2001	0510							2	0001	106 <
	W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
		CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,
		HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,	LS,	LT,
		LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,	RU,
		SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,
		YU,	ZA,	ZW,	AM,	AZ,	BY,	KG,	KΖ,	MD,	RU,	ТJ,	TM				
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,
		DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,	TR,	BF,
		ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	ΤG		
CA	2389	773			A1		2001	0510		CA 2	000-	2389	773		2	0001	106 <
EP	1237	849			A1		2002	0911		EP 2	000-	9730	61		2	0001	106
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,
		IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR						
JP	2003	5130	64		T		2003	0408		JP 2	001-	5347	58		2	0001	106
PRIORIT	Y APP	LN.	INFO	.:						GB 1	999-	2628	6		A 1	9991	105
										US 2	000-	2013	82P		P 2	0000	502
										WO 2	000-	GB42	49	,	W 2	0001	106
OTHER SO	OURCE	(S):			MAR	PAT	134:	3531	75								

The title compds. R4PZNR1R2 [I; R1, R2 = alkyl; R1R2 together form alkylene; Z = alkylene; P = a direct bond, X, Y, W, XY, YW, XYW (wherein W = O, S, NR3; R3 = H, alkyl; Y = UV; V = a direct bond, alkylene; U = CS, CO, SO2, C(:NR); R = H, OH, alkyl; X = O, NR6; R6 = H, alkyl, alkenyl, etc.); R4 = alkyl, alkenyl, alkynyl, etc.], useful in the activation of soluble guanylate cyclase, were prepared E.g., synthesis of the urea II, starting with 4-bromoaniline and 1-(3-aminopropyl)pyrrolidine, was given. Biol. data for compds. I (e.g., IC50 for inhibition of platelet aggregation) were presented.

IT 338980-58-8P 338980-88-4P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of amides and ureas as activators of soluble guanylate $\operatorname{cyclase}$)

RN 338980-58-8 ZCAPLUS

CN Benzamide, 2-[[(5-chloro-1-methyl-3-phenyl-1H-pyrazol-4-yl)methyl]thio]-N-[3-(dimethylamino)propyl]- (9CI) (CA INDEX NAME)

RN 338980-88-4 ZCAPLUS

CN Urea, N-[2-[[(5-chloro-1-methyl-3-phenyl-1H-pyrazol-4-yl)methyl]thio]phenyl]-N'-[3-(dimethylamino)propyl]- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 24 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:334328 ZCAPLUS Full-text

DOCUMENT NUMBER: 134:357554

TITLE: Application and formulation of isoxazole derivatives

as phosphodiesterase VII inhibitors

INVENTOR(S): Eggenweiler, Hans-Michael; Jonas, Rochus; Wolf,

Michael; Gassen, Michael; Welge, Thomas

PATENT ASSIGNEE(S): Merck Patent G.m.b.H., Germany

SOURCE: Ger. Offen., 6 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Fatent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA.	TENT	NO.			KIN	D	DATE			APPL	ICAT	ION 1	.00		D.	ATE		
	1995 2389															9991 0001		
WO	2001	0321	75		A1		2001	0510		WO 2	000-	EP10	239		2	0001	018	<
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		CZ,	DE,	DK,	DM,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,	
		IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	KΖ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MA,	
		MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	
		SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZW		
	RW:	,	,	,		,		,	,	SZ,	,	,	,			,	,	
		DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,	BF,	ΒJ,	
		,		,		,	,	,		MR,	,		,					
	2000																	
EP	1225	896			A1		2002	0731		EP 2	000-	9713	93		2	0001	018	
EP	1225																	
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,	
		ΙE,	SI,	LT,	LV,	FΙ,	RO,	MK,	CY,	AL								
	2002																	
	2003															0001		
	7816															0001	018	
	3009	45			Τ					AT 2			-		_	0001		
	2243				Т3					ES 2						0001		
	2002							0503		NO 2		_				0020		<
	6531				В1			0311		US 2					_	0020		
	2002							0910		MX 2						0020		
ZA	2002	0044	30		А		2003	0903		ZA 2	002-	4430			2	0020	603	

IN 2002KN00743 A 20050311 IN 2002-KN743 20020603
PRIORITY APPLN. INFO.: DE 1999-19953024 A 19991104
WO 2000-EP10239 W 20001018

OTHER SOURCE(S): MARPAT 134:357554

GΙ

$$\mathbb{R}^2$$
 \mathbb{R}^5 \mathbb{R}^4 \mathbb{R}^3

The invention concerns isoxazole derivs. of the formula (I) to be used as phosphodiesterase VII inhibitors and their drug formulations. In I R1,R2,R3,R4 = Hal,OA,SA,A,H,COOA,CN,CONA1A2,R5COOA1; A1,A2 = H, A, Alkenyl, Cycloalkyl, alkylene cycloalkyl; A = C1-C10 alkyl, Hal = F, C1, Br, J. Non-physiol. salts and solvates of the compds. can be used too. The compns. are used for the therapy of asthma, chronic bronchitis, dermatitis, autoimmune diseases etc. Thus an eye-drop solution contained 1 g of the formula I compound; further components in g were: NaH2PO4x2H2O 9.38; Na2HPO4x12H2O 28.48; benzalkonium chloride 0.1; double distilled water 940; pH 6.8.

IT 303995-75-7 303995-80-4 320424-92-8 338394-43-7 338402-64-5 338403-15-9

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses) (application and formulation of isoxazole derivs. as phosphodiesterase VII inhibitors)

RN 303995-75-7 ZCAPLUS

CN Benzoic acid, 4-[[2-(4-cyano-3-phenyl-5-isoxazolyl)ethenyl]amino]- (9CI) (CA INDEX NAME)

RN 303995-80-4 ZCAPLUS

CN 4-Isoxazolecarboxylic acid, 5-[2-[(4-carboxyphenyl)amino]ethenyl]-3-phenyl-, 4-methyl ester (9CI) (CA INDEX NAME)

RN 320424-92-8 ZCAPLUS

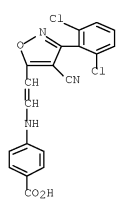
CN Benzoic acid, 4-[[2-[3-(2-chlorophenyl)-4-cyano-5-isoxazolyl]ethenyl]amino]- (9CI) (CA INDEX NAME)

RN 338394-43-7 ZCAPLUS

CN 4-Isoxazolecarboxylic acid, 5-[2-[(4-carboxyphenyl)amino]ethenyl]-3-(2,6-dichlorophenyl)-, 4-methyl ester (9CI) (CA INDEX NAME)

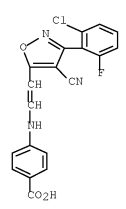
RN 338402-64-5 ZCAPLUS

CN Benzoic acid, 4-[[2-[4-cyano-3-(2,6-dichlorophenyl)-5-isoxazolyl]ethenyl]amino]- (9CI) (CA INDEX NAME)



RN 338403-15-9 ZCAPLUS

CN Benzoic acid, 4-[[2-[3-(2-chloro-6-fluorophenyl)-4-cyano-5-isoxazolyl]ethenyl]amino]- (9CI) (CA INDEX NAME)



L89 ANSWER 25 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:265369 ZCAPLUS Full-text

DOCUMENT NUMBER: 134:295620

TITLE: Preparation and effect of 4-methoxyphenylpropionic

acid derivatives useful in insulin resistance

improvement

INVENTOR(S): Shinoda, Masanobu; Emori, Eita; Matsuura, Fumiyoshi;

Kaneko, Toshihiko; Ohi, Norihito; Kasai, Shunji; Yoshitomi, Hideki; Yamazaki, Kazuto; Miyashita, Sadakazu; Hibara, Taro; Seiki, Hisashi; Clark,

Richard; Harada, Hitoshi

PATENT ASSIGNEE(S): Eisai Co., Ltd., Japan SOURCE: PCT Int. Appl., 350 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

GΙ

P.	ATENT	NO.			KIN	D	DATE			APPI	LICAT	ION	NO.		D	ATE		
WC	2001	 0251	81		A1	_	2001	0412		WO 2	2000-	JP67	88		2	0000	929	<
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	RW:	AT, PT,	•	CH,	CY,	DE	, DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	
TV	V 2621	85			В		2006	0921		TW 2	2000-	8912	0087		2	0000	928	
CA	2385	081			A1		2001	0412		CA 2	2000-	2385	081		2	0000	929	<
JA	J 2000	7449	9		Α		2001	0510		AU 2	2000-	7449	9		2	0000	929	<
JA	J 7762	67			В2		2004	0902										
E	2 1216	980			A1		2002	0626		EP 2	2000-	9629	93		2	0000	929	
	R:	•	BE, FI,	•	DE,	DK	, ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,	
NZ	3 5177	19			Α		2004	1029		NZ 2	2000-	5177	19		2	0000	929	
US	6884	821			В1		2005	0426		US 2	2002-	8891	6		2	0000	929	
PRIORIT	TY APP	LN.	INFO	.:						JP 1	L999-	2820	79		A 1	9991	001	
										JP 1	L999-	3694	42		A 1	9991	227	
										JP 2	2000-	3879	5		A 2	0000	216	
										JP 2	2000-	1042	60		A 2	0000	406	
										WO 2	2000-	JP67	88		W 2	0000	929	
OTHER S	SOURCE	(S):			MAR	PAT	134:	2956	20									

$$F_{3}C$$

$$NH$$

$$MeO$$

$$Me$$

$$Me$$

$$Me$$

$$I$$

AB Title compds. [Y:L:X:TZM:CWR1; R1 is hydrogen, hydroxyl, alkyl; L is single bond, double bond, alkylene; M is single bond, alkylene; T is single bond, alkylene; W is carboxyl, amide; X is oxygen, alkenylene; Y is aromatic hydrocarbon; Z is aromatic hydrocarbon; colon represents single, or double bond], salts, esters, and hydrates are prepared and are useful in prevention or treatment of diabetes and X-syndrome. Thus, the title compound I was prepared and biol. tested.

IT 334012-76-9P 334012-77-0P 334012-78-1P 334012-79-2P 334012-80-5P 334012-85-0P 334012-86-1P 334012-87-2P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation and effect of methoxyphenylpropionic acid derivs. useful in insulin resistance improvement as PPAR agonists)

RN 334012-76-9 ZCAPLUS

CN Benzenepropanoic acid, 4-methoxy- α -(1-methylethoxy)-3-[[[(1-methyl-3-phenyl-1H-pyrazol-5-yl)carbonyl]amino]methyl]- (9CI) (CA INDEX NAME)

RN 334012-77-0 ZCAPLUS

CN Benzenepropanoic acid, 4-methoxy- α -(1-methylethoxy)-3-[[[(1-methyl-5-phenyl-1H-pyrazol-3-yl)carbonyl]amino]methyl]- (9CI) (CA INDEX NAME)

RN 334012-78-1 ZCAPLUS

CN Benzenepropanoic acid, 4-methoxy- α -(1-methylethoxy)-3-[[[(5-phenyl-3-isoxazolyl)carbonyl]amino]methyl]- (9CI) (CA INDEX NAME)

RN 334012-79-2 ZCAPLUS

CN Benzenepropanoic acid, 4-methoxy- α -(1-methylethoxy)-3-[[[[1-methyl-5-(2-pyridinyl)-1H-pyrazol-3-yl]carbonyl]amino]methyl]- (9CI) (CA INDEX NAME)

RN 334012-80-5 ZCAPLUS

CN Benzenepropanoic acid, 4-methoxy- α -(1-methylethoxy)-3-[[[[1-methyl-3-(2-pyridinyl)-1H-pyrazol-5-yl]carbonyl]amino]methyl]- (9CI) (CA INDEX NAME)

RN 334012-85-0 ZCAPLUS

CN Benzenepropanoic acid, $3-[[[5-(2-\text{chloropheny1})-3-\text{isoxazolyl}]\text{carbonyl}]\text{amino}]\text{methyl}]-4-\text{methoxy}-\alpha-(1-\text{methylethoxy})-(9CI)$ (CA INDEX NAME)

RN 334012-86-1 ZCAPLUS

CN Benzenepropanoic acid, 4-methoxy- α -(1-methylethoxy)-3-[[[(3-methyl-1-phenyl-1H-pyrazol-5-yl)carbonyl]amino]methyl]- (9CI) (CA INDEX NAME)

RN 334012-87-2 ZCAPLUS

CN Benzenepropanoic acid, 4-methoxy- α -(1-methylethoxy)-3-[[[(5-methyl-1-phenyl-1H-pyrazol-3-yl)carbonyl]amino]methyl]- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 26 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:137199 ZCAPLUS Full-text

DOCUMENT NUMBER: 134:178561

TITLE: Preparation of heterocyclylmethyl substituted benzoic

acids for the treatment of diabetes mellitus

INVENTOR(S): Hargreaves, Rodney Brian; Whittamore, Paul Robert Owen

PATENT ASSIGNEE(S): AstraZeneca AB, Swed.; AstraZeneca AB

SOURCE: PCT Int. Appl., 59 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:	CENT :	NO.			KIN	D	DATE			APPL	ICAT	ION 1	NO.		D.	ATE		
WO	2001	 0126			A1	_	2001	0222		 WO 2	000-	 GB31	 26		2	0000	 814	<
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		CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	
		ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,	LS,	LT,	LU,	
		LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	
		SG,	SI,	SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZW
	RW:	GH,	GM,	KΕ,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,	
		DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	
		CF,	CG,	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,	ΤG				
CA	2381	090			A1		2001	0222	1	CA 2	000-	2381	090		2	0000	814	<
BR	2000	0133	74		Α		2002	0507		BR 2	000-	1337	4		2	0000	814	<
ΕP	1210	339		A1 20020605 EP 2000-953309 200										0000	814			
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	IE,	SI,	LT,	LV,	FΙ,	. RO,	MK,	CY, A	łΓ				
JP	200350737	72		Τ		2003	0225	JP	2	2001-517510		20000814	
AU	766790			В2		2003	1023	AU	J	2000-65823		20000814	
NZ	517060			Α		2003	1128	NZ	2 2	2000-517060		20000814	
ZA	200200067	70		Α		2003	0424	ZA	A 2	2002-670		20020124	
US	6787556			В1		2004	0907	US	3 2	2002-48392		20020129	
MX	2002PA015	597		Α		2002	0702	MX	ζ 2	2002-PA1597		20020214	
NO	200200076	54		Α		2002	0417	NO) 2	2002-764		20020215	<
PRIORITY	APPLN. 1	INFO.	. :					GB	3 :	1999-19413	Α	19990818	
								WO) 2	2000-GB3126	W	20000814	
000000	TIDOD (O)			1077	D 70 ETT	101	1705	C 1					

OTHER SOURCE(S): MARPAT 134:178561

GΙ

$$\mathbb{R}^2$$
 A $\mathbb{X} = \mathbb{Q}$ \mathbb{R}^3 \mathbb{Z} \mathbb{R}^3 $\mathbb{R}^$

The title compds. [I; Q, X, Y, Z = CRa, CRb:CRc, N (wherein Ra, Rb, Rc = H, halo, a bond, such that together with the nitrogen atom to which Y and Z are attached, they form a 5-6 membered aromatic ring); R1, R3 = alkyl, halo, haloalkyl, etc.; n = 0-2; A = alkylene, alkenylene, alkynylene optionally interposed by a heteroatom; R2 = (un)substituted aryl, heterocyclyl, cycloalkyl] which act as peroxisome proliferator activated receptor (PPAR) agonists, in particular states of insulin resistance including type 2 gamma receptors (PPAR) (data given), and so are useful therapeutically in the treatment of diabetes mellitus, were prepared E.g., a multi-step synthesis of the benzoic acid II was given.

IT 326912-92-9P 326912-93-0F 326912-94-1P

326912-98-5P 326912-99-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of heterocyclylmethyl substituted benzoic acids for the treatment of diabetes mellitus)

RN 326912-92-9 ZCAPLUS

CN Benzoic acid, 2-[[3-(3-bromo-4-methylphenyl)-1H-pyrazol-1-yl]methyl]-, ethyl ester (9CI) (CA INDEX NAME)

RN 326912-93-0 ZCAPLUS

CN Benzoic acid, 2-[[3-[3-bromo-4-(bromomethyl)phenyl]-1H-pyrazol-1-yl]methyl]-, ethyl ester (9CI) (CA INDEX NAME)

RN 326912-94-1 ZCAPLUS

CN Benzoic acid, 2-[[3-[3-bromo-4-[(methylamino)methyl]phenyl]-1H-pyrazol-1-yl]methyl]-, ethyl ester (9CI) (CA INDEX NAME)

RN 326912-98-5 ZCAPLUS

CN Benzoic acid, 2-[[3-(3-chloro-4-methylphenyl)-1H-pyrazol-1-yl]methyl]-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & & & \\ \hline & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

RN 326912-99-6 ZCAPLUS

CN Benzoic acid, 2-[[3-[4-(bromomethyl)-3-chlorophenyl]-1H-pyrazol-1-yl]methyl]-, methyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 27 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:101128 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 134:147599

TITLE: Preparation of 2-pyrazolin-5-ones as inhibitors of

serine/threonine and tyrosine kinase activity

INVENTOR(S): Moset, Marina M.; Berlanga, Jose Maria Castellano;

Fernandez, Isabel F.; Calderwood, David J.; Rafferty,

Paul; Arnold, Lee

PATENT ASSIGNEE(S): Basf Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 226 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Fatent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PA:	TENT	NO.			KINI	O	DATE		-	APPL	ICAT	ION :	NO.		D.	ATE		
		2001 2001									WO 2	000-	 US20	628		2	0000	728	<
			AE, CR, HU, LU, SD,	AG, CU, ID, LV,	AL, CZ, IL, MA, SG,	AM, DE, IN, MD,	AT, DK, IS, MG,	AU, DM, JP, MK,	AZ, DZ, KE, MN, TJ,	BA, EE, KG, MW,	ES, KP, MX,	FI, KR, MZ,	GB, KZ, NO,	GD, LC, NZ,	GE, LK, PL,	GH, LR, PT,	GM, LS, RO,	HR, LT, RU,	
		RW:	DE,	DK,	ES,	FI,	FR,	GB,	SD, GR, GW,	IE,	IT,	LU,	MC,	NL,	PT,				
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	CA	2380																	<
		2000																	
	ΕP	1218	373			A2		2002	0703		EP 2	000-	9508	52		2	0000	728	
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	TR	2002							0923			002-	928			2	0000	728	
		2003																	
		2004															0000		
	HU	2004	0054	0		А3		2004	0928										
		5168						2004	0924		NZ 2	000-	5168	50		2	0000	728	
	IN	2002	MN00	057		А		2006	0505		IN 2	002-	MN57			2	0020	116	
	ZA	2002	0004	77		А		2003	0422		ZA 2	002-	477			2	0020	118	
		2002						2002	0312			002-					0020	130	<
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	ВG	1063	92			Α		2002	1229		BG 2	002-	1063	92		2	0020	206	
PRIOF	RIT	Y APP	LN.	INFO	.:							999- 000-							
OTHER		OLID OLI	(0)			MADI	~ T	104.	1 475										

OTHER SOURCE(S): MARPAT 134:147599

The title compds. [I; R = (un)substituted alkyl, aryl, cycloalkyl, etc.; R1 = H, AZ; R2 = H, (un)substituted alkyl, aryl, etc.; A = (CH2)n, (CH2)nNH, (CH2)nO, etc.; Z = H, alkyl, aralkyl, etc.] which are inhibitors of serine/threonine and tyrosine kinase activity, were prepared and formulated. Thus, reacting 3-cyclopropyl-2-pyrazolin-5-one with 4,5-dimethylpyrrole-2-carboxaldehyde in the presence of piperidine in EtOH afforded 30% I [R = 4,5-dimethylpyrrol-2-yl; R1 = cyclopropyl]. All exemplified compds. I inhibit KDR kinase at 50 μ M and some of them also significantly inhibit other PTKs such as lck at \leq 50 μ M, and cdc2 at < 50 μ M. Several of the tyrosine kinases, whose activity is inhibited by the compds. I are involved in angiogenic processes. Thus, the compds. I can ameliorate disease states where angiogenesis or endothelial cell hyperproliferation is a factor. These compds. I can be used to treat cancer and hyperproliferative disorders.

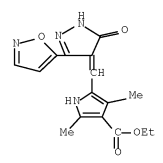
IT 324549-32-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of 2-pyrazolin-5-ones as inhibitors of serine/threonine and tyrosine kinase activity)

RN 324549-32-8 ZCAPLUS

CN 1H-Pyrrole-3-carboxylic acid, 5-[[1,5-dihydro-3-(5-isoxazolyl)-5-oxo-4H-pyrazol-4-ylidene]methyl]-2,4-dimethyl-, ethyl ester (9CI) (CA INDEX NAME)



L89 ANSWER 28 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:31489 ZCAPLUS Full-text

DOCUMENT NUMBER: 134:100865

TITLE: Preparation of 1-(4-quinoly1)-1H-pyrazoles as

agrochemical fungicides

INVENTOR(S): Emeric, Gilbert; Gary, Stephanie; Gerusz, Vincent;

Gourlaouen, Nelly; Hartmann, Benoit; Huser, Nathalie; Lachaise, Helene; Le Hir De Fallois, Loic; Perez,

Joseph; Wegmann, Thomas

PATENT ASSIGNEE(S): Aventis CropScience SA, Fr. SOURCE: PCT Int. Appl., 267 pp.

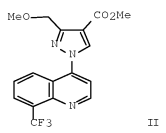
CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

P.	ATENT	NO.			KIN	D	DATE			APPL	ICAT	ION 1	ΝΟ.		D.	ATE	
M	0 2001	0023	85		A1		2001	0111	,	WO 2	000-	FR18	16		2	0000	629 <
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		CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,
		ID,	IL,	IN,	IS,	JP,	KΕ,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,	LS,	LT,	LU,
		LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NΖ,	PL,	PT,	RO,	RU,	SD,	SE,
		SG,	SI,	SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,	YU,	ZA,
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	RW:	GH,	GM,	KΕ,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,
		DE,	DK,	ES,	FΙ,	FR,	GB,	GR,	IE,	ΙT,	LU,	MC,	NL,	PT,	SE,	BF,	BJ,
		CF,	CG,	CI,	CM,	GΑ,	GN,	GW,	${ m ML}$,	MR,	ΝE,	SN,	TD,	ΤG			
F1	R 2795	726			A1		2001	0105		FR 1	999-	8596			1	9990	630 <
PRIORI'	TY APP	LN.	INFO	.:						FR 1	999-	8596			A 1	9990	630
OTHER :	SOURCE	(S):			MAR:	PAT	134:	1008	65								



GΙ

AB R1R2 [I; R1 = (un)substituted 4-quinolyl; R2 = di- or trisubstituted pyrazolo] were prepared Thus, MeOCH2COCH2CO2Me was condensed with HC(OMe)2NMe2 and the product cyclocondensed with H2NNH2 to give Me 5-methoxymethylpyrazole-4-carboxylate which was N-arylated by 4-chloro-8-trifluoromethylquinoline to give title compound II. Data for biol. activity of I were given.

IT 318492-52-3P 318492-66-9P

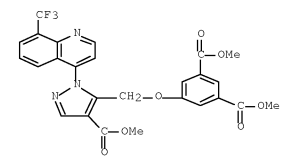
RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of 1-(4-quinolyl)-1H-pyrazoles as agrochem. fungicides)

RN 318492-52-3 ZCAPLUS

CN 1,3-Benzenedicarboxylic acid, 5-[[1-(7-chloro-4-quinolinyl)-4-(methoxycarbonyl)-1H-pyrazol-5-yl]methoxy]-, dimethyl ester (9CI) (CA INDEX NAME)

RN 318492-66-9 ZCAPLUS

CN 1,3-Benzenedicarboxylic acid, 5-[[4-(methoxycarbonyl)-1-[8-(trifluoromethyl)-4-quinolinyl]-1H-pyrazol-5-yl]methoxy]-, dimethyl ester (9CI) (CA INDEX NAME)



REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 29 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:5870 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 134:127200

TITLE: Activity of the new BASF strobilurin fungicide, BAS

500 F, against Plasmopara viticola on grapes

AUTHOR(S): Stierl, R.; Scherer, M.; Schrof, W.; Butterfield, E.

J.

CORPORATE SOURCE: Agricultural Center, BASF AG, Limburgerhof, 67114,

Germany

SOURCE: BCPC Conference--Pests & Diseases (2000),

(Vol. 1), 261-266 CODEN: BCDCAE

PUBLISHER: British Crop Protection Council

DOCUMENT TYPE: Journal LANGUAGE: English

AB BAS 500 F is the new, broad-spectrum strobilurin fungicide being developed by BASF. The compound provides excellent control of Plasmopara viticola, the pathogen which causes downy mildew of grapevines. Field trials, under practical conditions, have shown that BAS 500 F controls this disease effectively on leaves and berries. Microscopic studies revealed that this good control is due to high activity of the compound against several developmental stages of the pathogen. The zoospores are extremely sensitive

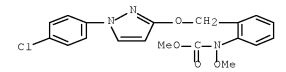
to BAS $500~\mathrm{F}$ and react to contact with lysis. If zoospores escape lysis, the germination of encysted zoospores is stopped effectively by a preventative treatment. After curative application, the compound stops further development of the mycelium in the leaves.

IT 175013-18-0, BAS 500F

RL: AGR (Agricultural use); BIOL (Biological study); USES (Uses) (activity against Plasmopara viticola on grapes)

RN 175013-18-0 ZCAPLUS

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)



REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 30 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:1406 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 134:127186

TITLE: BAS 500 F - the new broad-spectrum strobilurin

fungicide

AUTHOR(S): Ammermann, E.; Lorenz, G.; Schelberger, K.; Mueller,

B.; Kirstgen, R.; Sauter, H.

CORPORATE SOURCE: Agricultural Center, BASF AG, Limburgerhof, 67 114,

Germany

SOURCE: BCPC Conference--Pests & Diseases (2000),

(Vol. 2), 541-548 CODEN: BCDCAE

PUBLISHER: British Crop Protection Council

DOCUMENT TYPE: Journal LANGUAGE: English

BAS 500 F is the code number of the new, broad-spectrum strobilurin fungicide developed by BASF. As a foliar spray, it controls the major plant pathogens from the Ascomycete, Basidiomycete, Deuteromycete and Oomycete classes of fungi. BAS 500 F has protectant, curative, translaminar and locosystemic properties, and thus a broad and flexible application window. It is a highly active fungicide for cereals, peanuts and other field crops, grapes, vegetables, bananas, citrus and turf with excellent crop safety. The expected dose rate ranges from 50 - 250 g a.i./ha for food crops and from 280 - 560 g a.i./ha for turf. The compound has a favorable toxicol. and ecotoxicol. profile and is safe to users and the environment. It is classified by US-EPA as a "reduced risk candidate". BAS 500 F is being developed and registered as a solo product and with various premix partners, in a range of formulations. Market introduction is expected for the 2002 season.

IT 175013-18-0, BAS 500F

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study); USES (Uses)

(broad-spectrum strobilurin fungicide)

RN 175013-18-0 ZCAPLUS

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 31 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:1386 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 134:127181

TITLE: Activity of the new BASF strobilurin fungicide, BAS

500 F, against Septoria tritici on wheat

AUTHOR(S): Stierl, R.; Merk, M.; Schrof, W.; Butterfield, E. J. CORPORATE SOURCE: Agricultural Center, BASF AG, Limburgerhof, 67 114,

Germany

SOURCE: BCPC Conference--Pests & Diseases (2000),

(Vol. 3), 859-864 CODEN: BCDCAE

PUBLISHER: British Crop Protection Council

DOCUMENT TYPE: Journal LANGUAGE: English

AB BAS 500 F is the new, broad-spectrum strobilurin fungicide being developed by BASF. Field trials under practical conditions have shown that BAS 500 F effectively controls Septoria tritici blotch of wheat resulting in an increased yield in comparison to other strobilurin and triazole fungicides. Glasshouse and semifield trials in combination with microscopic techniques, i.e. conventional fluorescent and confocal laser scanning microscopic techniques, revealed that this good control is due to a very high intrinsic activity of the compound against several development stages of the pathogen. After a preventative treatment, germination of pycnidiospores is effectively stopped by BAS 500 F. Under curative conditions, the compound stops further development of the mycelium in the leaves and the subsequent yellowing and necrosis of leaf tissue.

IT 175013-18-0, BAS 500F

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study); USES (Uses)

(activity against Septoria tritici on wheat)

RN 175013-18-0 ZCAPLUS

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 32 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2000:900389 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 134:38252

TITLE: Synergistic fungicidal combinations of benzophenones

with strobilurins, cyanoimidazoles, and carbonic acid

amides

INVENTOR(S): Dalton, Ian Paul

PATENT ASSIGNEE(S): Novartis Ag, Switz.; Novartis-Erfindungen

Verwaltungsgesellschaft M.B.H.

PCT Int. Appl., 25 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.							DATE					ION				ATE		
		2000				 A1		2000			 WO 2		 EP54				0000	613	<
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			ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,	LS,	LT,	LU,	
			LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	
			SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UΖ,	VN,	YU,	
			ZA,	ZW,	ΑM,	AΖ,	BY,	KG,	KΖ,	MD,	RU,	ΤJ,	TM						
		RW:						${ m MZ}$,											
			DE,	DK,	ES,	FΙ,	FR,	GB,	GR,	ΙE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	
			CF,	CG,	CI,	CM,		GN,											
		1185				A1		2002			EP 2	000-	9512	83		2	0000	613	<
	EP	1185				В1		2003											
		R:						ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,	
						LV,													
		2000				Α		2002				000-					0000		<
		2003				Τ		2003				001-				_	0000		
		2412				Τ		2003	0615			000-					0000		
		1185				I		2003 2003 2004 2002	1031			000-					0000		
		2200				T3		2004	0316			000-					0000		
		2002		46		A1		2002	0808		US 2	001-	9976	0.7		2	0011	129	
		6689				В2		2004	0210					_					
PRIC	RIT	Y APP	LN.	INFO	.:							999-					9990		
												999-					9990		
												999-					9990		
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												999- 999-					9990 9990		
											GD I	シンツー	1002	U		a I	シンプU	014	

GB	1999-13822	Α	19990614
GB	1999-13824	Α	19990614
GB	1999-13826	Α	19990614
GB	1999-13827	Α	19990614
WO	2000-EP5433	W	20000613

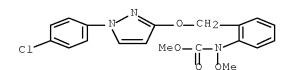
GΙ

$$R^1$$
 R^2 R^3 R^2 R^3

The invention relates to a method of combating phytopathogenic diseases on crop plants which comprises applying to the crop plants or the locus thereof being infested with said phytopathogenic disease an effective amount of a combination of a benzophenone I (R1 = methoxy, Me; R2 = C1-C4alkoxy, 2-halogenbenzyloxy; R3 = C1-C4alkoxy; R4 = C1-C4alkyl, halo, or trifluoromethyl; R5 = H, halo, C1-C4alkoxy, trifluoromethyl, or nitro) in association with a compound selected from strobilurins, cyanoimidazoles, and carbonic acid amides.

RN 175013-18-0 ZCAPLUS

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)



REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 33 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2000:367993 ZCAPLUS Full-text

DOCUMENT NUMBER: 133:1743

TITLE: Synergistic fungicidal mixtures

INVENTOR(S): Schelberger, Klaus; Scherer, Maria; Saur, Reinhold;

Sauter, Hubert; Muller, Bernd; Birner, Erich; Leyendecker, Joachim; Ammermann, Eberhard; Lorenz,

Gisela; Strathmann, Siegfried

PATENT ASSIGNEE(S): BASF A.-G., Germany SOURCE: PCT Int. Appl., 22 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PAT	TENT	NO.								APP1	LICAT	ION :	NO.		D.	ATE		
CA BR	W:	AE, CZ, IN, MD, SK, GH, DK, CG, 819	AL, DE, IS, MG, SL, GM, ES, CI,	AM, DK, JP, MK, TJ, KE, FI, CM,	AT, DM, KE, MN, TM, LS, FR, GA, A1	AU, EE, KG, MW, TR, MW, GB,	AZ, ES, KP, MX, TT, SD, GR, GW, 2000	BA, FI, KR, NO, TZ, SL, IE, ML, 0602	BB, GB, KZ, NZ, UA, SZ, IT, MR,	BG, GD, LC, PL, UG, TZ, LU, NE, CA	1999- BR, GE, LK, PT, US, UG, MC, SN, 1999-	BY, GH, LR, RO, UZ, ZW, NL, TD, 2351	CA, GM, LS, RU, VN, AT, PT, TG 819	CH, HR, LT, SD, YU, BE,	CN, HU, LU, SE, ZA, CH, BF,	ID, LV, SG, ZW CY,	CU, IL, MA, SI, DE, CF,	< <
EP	1130				В1		2003											
	R:				DE, LV,		•	FR,	GB,	GR,	, IT,	LI,	LU,	NL,	SE,	MC,	PT,	
HU	2001	0441	4		A2		2002	0328		HU 2	2001-	4414			1	9991	106	<
JP	2002	5303			Τ		2002	0917			2000-				1	9991	106	
AT	2453	54			T		2003	0815			1999-				1	9991	106	
NZ	5121	91			Α		2003	0829		NZ :	1999-	5121	91		1	9991	106	
AU	7675	77			В2		2003	1120			2000-				1	9991	106	
PT	1130	967			T		2003	1128			1999-				1	9991	106	
CZ	2934	37			В6		2004	0414		CZ 2	2001-	1737			1	9991	106	
ES	2204	196			Т3		2004	0416		ES 3	1999-	9724	95		1	9991	106	
RU	2244	420			C2		2005	0120		RU 2	2001-	1165	95		1	9991	106	
SK	2847	47			В6		2005	1103		SK 2	2001-	678			1	9991	106	
IL	1431	01			A		2005	1218		IL 3	1999-	1431	01		1	9991	106	
MX	2001	PA04	959		A		2001	0731		MX 2	2001-	PA49	59		2	0010	517	<
US	6503	936			В1		2003	0107		US 2	2001-	8560	34		2	0010	517	
BG	1055	37			A		2002	0430		BG 2	2001-	1055	37		2	0010	522	<
BG	6500	3			В1		2006	1229										
ZA	2001	0049	62		A		2002	0618		ZA 2	2001-	4962			2	0010	618	
IN	2001	CN00	836		А		2005	0304		IN 2	2001-0	CN83	6		2	0010	618	
RIORIT	Y APP	LN.	INFO	.:							1998-				A 1	9981	119	
										WO :	1999-:	EP85	12		W 1	9991	106	
THER SO	DURCE	(S):			MARI	PAT	133:	1743										

AB A synergistic fungicidal mixture contains a carbamate I [X = CH or N; n = 0, 1 or 2; R = halo or C1-4 (halo)alkyl] and a copper compound

IT 216659-76-6 271249-36-6

RL: AGR (Agricultural use); BIOL (Biological study); USES (Uses) (synergistic fungicidal mixture)

RN 216659-76-6 ZCAPLUS

CN Carbamic acid, [2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester, mixt. with copper hydroxide sulfate (Cu4(OH)6(SO4)) (9CI) (CA INDEX NAME)

CM 1

CRN 175013-18-0 CMF C19 H18 C1 N3 O4

CM 2

CRN 1333-22-8

CMF Cu . H O . O4 S

CCI TIS

CM 3

CRN 14808-79-8

CMF 04 S

CM 4

CRN 14280-30-9

CMF H O

он-

CM 5

CRN 7440-50-8

CMF Cu

Cu

RN 271249-36-6 ZCAPLUS

CN Carbamic acid, methoxy[2-[[[1-(4-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-, methyl ester, mixt. with copper hydroxide sulfate (Cu4(OH)6(SO4)) (9CI) (CA INDEX NAME)

CM 1

CRN 175013-22-6 CMF C20 H21 N3 O4

CM 2

CRN 1333-22-8

CMF Cu . H O . O4 S

CCI TIS

CM 3

CRN 14808-79-8

CMF 04 S

CM 4

CRN 14280-30-9

CMF H O

он-

CM 5

CRN 7440-50-8

CMF Cu

Cu

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 34 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2000:349202 ZCAPLUS Full-text

DOCUMENT NUMBER: 132:344443

TITLE: Synergistic fungicidal compositions.

INVENTOR(S): Mauler-Machnik, Astrid; Wachendorf-Neumann, Ulrike;

Gayer, Herbert

PATENT ASSIGNEE(S): Bayer A.-G., Germany SOURCE: Ger. Offen., 18 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:	TENT	NO.			KIN		DATE			APPL	ICAT	ION :	NO.		D.	ATE	
	1993				A1			0525		DE 1						9990	823 <
ΙN	1999	BO00	745		А		2005	0304		IN 1						9991	102
	2351	500			A1		2000	0602		CA 1	999-	2351	500		1	9991	108 <
WO	2000	0304	40		A2		2000	0602		WO 1	999-	EP85	58		1	9991	108 <
WO	2000	0304	40		А3		2000	0831									
	W:	ΑE,	AL,	AM,	ΑT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CR,	CU,
		CZ,	DE,	DK,	DM,	EE,	ES,	FΙ,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,
		IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MA,
		MD,	MG,	MK,	MN,	MW,	MX,	NO,	NΖ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,
		SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UΖ,	VN,	YU,	ZA,	ZW	
	RW:									TZ,							DE,
		DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,
										NE,							
AU	2000	1046	0		Α		2000	0613		AU 2	000-	1046	0		1	9991	108 <
	7524																
BR	9915	518			Α		2001	0717		BR 1	999-	1551	8		1	9991	108 <
EP	1130	963			A2		2001	0912		EP 1	999-	9539	75		1	9991	108 <
EP	1130	963			В1		2005	0302									
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FΙ,	RO										
TR	2001	0137	9		T2		2001	1121		TR 2	001-	2001	0137	9	1	9991	108 <
	2001						2002	0328		HU 2	001-	4483			1	9991	108 <
TR	2001	0381	0		T2		2002	0621		TR 2	001-	2001	0381	0	1	9991	108
TR	2001	0381	1		Т2		2002	0621		TR 2	001-	2001	0381	1	1	9991	108
JP	2002	5302														9991	108
EP	1506	711			A2		2005	0216		EP 2	004-	2446	3		1	9991	108
EP	1506	711			АЗ												
	R:		BE, FI,		DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,

AT 28	89750	T	20050315	ΑT	1999-953975		19991108	
PT 13	130963	T	20050630	PT	1999-953975		19991108	
ES 22	238853	T3	20050901	ES	1999-953975		19991108	
TW 52	21994	В	20030301	${\tt TW}$	1999-88119807		19991115	
US 6	559136	B1	20030506	US	2001-856023		20010516	
MX 20	001PA05029	A	20000827	${\rm MX}$	2001-PA5029		20010518	<
US 20	003161896	A1	20030828	US	2003-371770		20030221	
PRIORITY A	APPLN. INFO.:			DE	1998-19853559	A1	19981120	
				DE	1999-19939841	Α	19990823	
				ΕP	1999-953975	A3	19991108	
				WO	1999-EP8558	W	19991108	
				US	2001-856023	A3	20010516	

OTHER SOURCE(S): MARPAT 132:344443

GΙ

AB The title compns. comprise the pyrimidine derivs. I [Z = (un)substituted Ph; X = halo; A = heterocyclyl, CO2Me or CHNHMe] and any of a large number of known fungicides.

IT 175013-18-0D, mixts. with pyrimidine derivs.

RL: AGR (Agricultural use); BIOL (Biological study); USES (Uses) (synergistic fungicidal compns.)

RN 175013-18-0 ZCAPLUS

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

L89 ANSWER 35 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2000:335180 ZCAPLUS Full-text

DOCUMENT NUMBER: 132:330856

TITLE: Synergistic fungicidal combinations comprising a

thieno[2,3-d]pyrimidin-4-one

INVENTOR(S): Walter, Harald; Forster, Birgit; Knauf-beiter,

Gertrude

PATENT ASSIGNEE(S): Novartis Ag, Switz.; Novartis-Erfindungen

Verwaltungsgesellschaft Mbh

SOURCE: PCT Int. Appl., 44 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	TENT	NO.			KIN	D										ATE		
WO	2000	0272	00		A1	_	2000				 999-					9991	104	<
	W:	ΑE,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	ВG,	BR,	BY,	CA,	CH,	CN,	CR,	CU,	
		CZ,	DE,	DK,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	
		IS,	JP,	KΕ,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	
		MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	
		SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZW			
	RW:	GH,	GM,	KΕ,	LS,	MW,	SD,	SL,	SZ,	ΤZ,	UG,	ZW,	AT,	BE,	CH,	CY,	DE,	
		DK,	ES,	FΙ,	FR,	GB,	GR,	ΙE,	ΙT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	
							GW,											
	2347																	
	9915																	
	2001																	
EP	1124																	
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,	
		,	,	,	LV,	,												
	2001																	<
	2002										000-				_			
	7562						2003				000-		-		_	9991		
	2228						2002				999-					9991		
	2001										001-							<
	2001						2005				001-		-			0010		
	2002				A1		2002	0321			001-					0010		<
PRIORIT	Y APP	LN.	INFO	.:							998-							
										WO 1	999-	EP84	49	,	W 1	9991	104	
OTHER S(GI	OURCE	(S):			MAR	PAT	132:	3308	56									

$$\begin{array}{c|c} & \circ & \\ &$$

The title compns. comprise a thieno[2,3-d]pyrimidin-4-one I (R1= halo; R2, R3 = C2-5 alkyl or methylcyclopropyl) in association with either an azole fungicide, an anilinopyrimidine fungicide, a morpholine fungicide, or strobilurin compound, a pyrrole derivative, a phenylamide, a dithiocarbamate fungicide (mancozeb, maneb, metiram or zineb), a copper compound (copper hydroxide, copper oxychloride, copper sulfate or oxine-copper), sulfur, prochloraz, triflumizole, pyrifenox, acibenzolar-S-Me, chlorothalonil, cymoxanil, dimethomorph, famoxadone, quinoxyfen, fenpropidine, spiroxamine, triazoxide, BAS 50001F, hymexazole, pecycuron, fenamidone, MON65500, or quazatine.

IT 175013-33-9D, BAS 50001F, mixts. with thieno[2,3-d]pyrimidin-4-one derivs.

RL: AGR (Agricultural use); BIOL (Biological study); USES (Uses) (synergistic fungicidal compns.)

RN 175013-33-9 ZCAPLUS

CN Carbamic acid, [2-[[[1-(4-fluorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 36 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2000:235080 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 133:4618

TITLE: Novel retinoic acid receptor α agonists:

syntheses and evaluation of pyrazole derivatives

AUTHOR(S): Kikuchi, Kouichi; Hibi, Shigeki; Yoshimura, Hiroyuki;

Tai, Kenji; Hida, Takayuki; Tokuhara, Naoki; Yamauchi,

Toshihiko; Nagai, Mitsuo

CORPORATE SOURCE: Tsukuba Basic Research Laboratories for Drug

Discovery, Eisai Co. Ltd., Tsukuba, 300-2635, Japan

SOURCE: Bioorganic & Medicinal Chemistry Letters (2000

Ι

), 10(7), 619-622

CODEN: BMCLE8; ISSN: 0960-894X

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

GΙ

TΤ

AB A series of pyrazole derivs. have been prepared as retinoic acid receptor (RAR) agonists. One of them, 4-[5-(1,5-diisopropyl-1H-3-pyrazolyl)-1H-2-pyrrolyl] benzoic acid (I), which possesses a 2,5-disubstituted pyrrole moiety, showed selective transactivation activity for the RAR α receptor, and had highly potent cell-differentiating activity on HL-60 cells.

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn, biol. activity and structure activity relationships of (alkylisopropylpyrazolyl)pyrrolylbenzoic acids as retinoic acid

receptor α agonists)

RN 270585-16-5 ZCAPLUS

270585-16-5P

CN Benzoic acid, 4-[4-[1-(2,5-dimethylphenyl)-5-(1-methylethyl)-1H-pyrazol-3-

yl]-1,4-dioxobutyl]-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{Me} \\ \text{N} \\ \text{i-Pr} \end{array} \qquad \begin{array}{c} \text{O} \\ \text{CH}_2 - \text{CH}_2 - \text{CH}_2 \end{array} \qquad \begin{array}{c} \text{O} \\ \text{C} \\ \text{OMe} \end{array}$$

REFERENCE COUNT: 31 THERE ARE 31 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 37 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2000:123270 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 132:151816

TITLE: Preparation of sulfonylureidopyrazoles as endothelin

converting enzyme inhibitors

INVENTOR(S): Hasegawa, Hirohiko; Yamazaki, Kazuto; Kanaoka, Shoji;

Ohashi, Naohito

PATENT ASSIGNEE(S): Sumitomo Pharmaceuticals Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 54 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Fatent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

GI

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2000053649	A	20000222	JP 1998-226684	19980811 <
PRIORITY APPLN. INFO.:			JP 1998-226684	19980811
OTHER SOURCE(S):	MARPAT	132:151816		

- AB The title compds. I [R1 = alkyl, etc.; R2, R3 = H, alkyl, etc.; R4 = H, halo, etc.; R5 = H, alkyl, etc.; R6 = RB1YA1; A1, B1 = alkylene, etc.; Y = OCO, etc.; R = H, cycloalkyl, etc.] are prepared I are useful in the treatment of cardiovascular diseases such as hypertension, arteriosclerosis, myocardial infarction, etc., cerebrovascular diseases, kidney diseases, asthma, complications of diabetes, endotoxin shock, etc. 4-Cyano-1-phenyl-3-benzyloxycarbonylmethyl-5-[3-(4-chlorobenzenesulfonyl)ureido]-(1H)pyrazole in vitro showed IC50 of 0.058 μ M against endothelin converting enzyme.
- IT 257954-72-6P 257954-77-1P 257954-82-8P
 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

(Reactant or reagent)

(preparation of sulfonylureidopyrazoles as endothelin converting enzyme inhibitors)

RN 257954-72-6 ZCAPLUS

CN Benzoic acid, 4-[[(5-amino-4-cyano-1-phenyl-1H-pyrazol-3-yl)acetyl]amino]-, ethyl ester (9CI) (CA INDEX NAME)

RN 257954-77-1 ZCAPLUS

CN Benzoic acid, 3-[[(5-amino-4-cyano-1-phenyl-1H-pyrazol-3-yl)acetyl]amino]-, ethyl ester (9CI) (CA INDEX NAME)

Ph
$$CH_2$$
 CH_2 CH_2

RN 257954-82-8 ZCAPLUS

CN Benzoic acid, 2-[[(5-amino-4-cyano-1-phenyl-1H-pyrazol-3-yl)acetyl]amino]-, ethyl ester (9CI) (CA INDEX NAME)

L89 ANSWER 38 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2000:3379 ZCAPLUS Full-text

DOCUMENT NUMBER: 132:35697

TITLE: Preparation and fungicidal activity of pyrazole

derivatives

INVENTOR(S): Desbordes, Philippe; Ellwood, Charles; Perez, Joseph;

Vors, Jean Pierre

PATENT ASSIGNEE(S): Rhone Poulenc Agrochimie, Fr.

SOURCE: Fr. Demande, 54 pp.

CODEN: FRXXBL

DOCUMENT TYPE: Fatent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PA	TENT	NO.			KIN	D	DATE			APPL	ICAT	ION :	NO.		D.	ATE	
	2773						1999			 FR 1	 997-	 1683	5 5		1		229 <
	2773						2000										
WO	9933	812			A1		1999	0708		WO 1	998-	FR28	42		1	9981	223 <
	W:	AL,	ΑM,	ΑT,	ΑU,	ΑZ,	ΒA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,	DE,
		DK,	EE,	ES,	FΙ,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,
		KE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,
		MW.	MX,	NO.	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG.	SI,	SK,	SL,	TJ,	TM,
		•	•	•	•	•	UZ,	•	•	•	•	•	•	•	•	•	•
		ŢJ,	TM	,	,	,	,	•	•	,	,	,	,	,	,	,	•
	RW:	GH,		KE.	LS.	MW.	SD.	S7.	UG.	7W.	AT.	BE.	CH.	CY.	DE.	DK.	ES.
		•			•		IT,	•			•						
				•			MR,			•		<i>0</i> _,	21,	20,	01,	00,	01,
7.11	9918		•	•	•	•	1999	•	•	•		1 2 2 1	a		1	9991	223 <
	9811				А		1999	1102									229 <
PRIORIT	Y APP	LN.	TNF.O	.:							997–		-		A 1		
										WO 1	998-	FR28	42		W 1	9981	223
OTHER S	OURCE	(S):			MAR:	PAT	132:	3569	7								

The title compds. I [G = R50Q1:CMeC(:Q2)R4, R5SQ1:CMeC(:Q2)R4, R6CH:CMeC(:Q2)R4, etc.; Q1 = N, CN, Q2 = O, S; Z = H, alkyl, haloalkyl, etc.; W = bond, O, S, SO, SO2, etc.; X1, X2, X3 = H, halo, OH, NO2, etc.; X4 = H, halo, alkyl, etc.; R3 = H, halo, alkyl, haloalkyl, etc.], possessing fungicidal activity, were prepared E.g., Me (E)-2-[2-[(4-methoxycarbonyl-1-methyl-5-phenoxy-1H-pyrazol-3-yl)oxymethyl]phenyl]-3- methoxyacrylate was prepared Fungicidal activity of I was tested against Plasmapora viticola, Puccinia recondita, Septoria tritici, etc.

IT 252280-47-0P 252280-48-1F 252280-49-2F

252280-50-5F 252280-51-6F 252280-52-7F

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation and fungicidal activity of pyrazole derivs.)

RN 252280-47-0 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[[(1-methyl-5-phenyl-1H-pyrazol-3-yl)oxy]methyl]-, methyl ester, (α E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 252280-48-1 ZCAPLUS

CN Benzeneacetic acid, α -(methoxyimino)-2-[[(5-phenyl-1H-pyrazol-3-yl)oxy]methyl]-, methyl ester, (α E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 252280-49-2 ZCAPLUS

CN Benzeneacetamide, α -(methoxyimino)-N-methyl-2-[[(5-phenyl-1H-pyrazol-3-yl)oxy]methyl]-, (α E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 252280-50-5 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[(5-phenyl-1H-pyrazol-3-yl)oxy]methyl]-, methyl ester, (α E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 252280-51-6 ZCAPLUS

CN Carbamic acid, methoxy[2-[[(5-phenyl-1H-pyrazol-3-yl)oxy]methyl]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 252280-52-7 ZCAPLUS

CN Benzeneacetic acid, α -methoxy-2-[[(5-phenyl-1H-pyrazol-3-yl)oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

L89 ANSWER 39 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1999:813423 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 132:20080

TITLE: Synergistic fungicidal compositions comprising a

strobilurine analog and a phosphite

INVENTOR(S):
Duvert, Patrice

PATENT ASSIGNEE(S): Rhone Poulenc Agro S. A., Fr.

SOURCE: Fr. Demande, 19 pp.

CODEN: FRXXBL

DOCUMENT TYPE: Fatent
LANGUAGE: French
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
FR 2778314	A1	19991112	FR 1998-6052	19980507 <
FR 2778314	В1	20020614		
PRIORITY APPLN. INFO.:			FR 1998-6052	19980507

OTHER SOURCE(S):

MARPAT 132:20080

GΙ

$$\begin{array}{c} \text{CH2-O} \\ \text{N} \end{array} \begin{array}{c} \text{T} \\ \text{N} \end{array} \begin{array}{c} \text{R} \\ \text{N} \end{array}$$

AB Synergistic fungicidal compns. comprising a strobilurine analog I [T = CH or N; R = H, halo or (halo)alkyl; n = 0, 1-5] and a phosphite, such as fosetyl-Al.

IT 251636-76-7

RL: AGR (Agricultural use); BIOL (Biological study); USES (Uses) (synergistic fungicidal composition)

RN 251636-76-7 ZCAPLUS

CN Carbamic acid, [3-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester, mixt. with aluminum tris(ethyl phosphonate) (9CI) (CA INDEX NAME)

CM 1

CRN 251636-75-6 CMF C19 H18 C1 N3 O4

CM 2

CRN 39148-24-8

CMF C2 H7 O3 P . 1/3 Al

●1/3 Al

ACCESSION NUMBER: 1999:722844 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 131:318921

TITLE: Synergistic fungicidal mixtures

INVENTOR(S): Schelberger, Klaus; Scherer, Maria; Saur, Reinhold; Eicken, Karl; Sauter, Hubert; Ammermann, Eberhard;

Grote, Thomas; Lorenz, Gisela; Strathmann, Siegfried

PATENT ASSIGNEE(S): BASF A.-G., Germany SOURCE: PCT Int. Appl., 23 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Fatent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	ENT :	NO.			KINI)	DATE			APPL	ICAT	ION :	NO.		D.	ATE		
WO		AL, LT, AM, AT,	AU, LV, AZ, BE,	BG, MK, KG,	BR, MX, MD,	BY, NO, TJ,	CA, NZ, TM	CN, PL,	CZ, RO,	GE, RU,	999- HU, SG,	ID, SI,	IL, SK,	IN, TR,	JP, UA,	KR, US,	KZ, ZA,	
CA	2330	PT, 607			A1		1999	1111		CA 1	999-	2330	607		1	9990	423	<
	9938						1999	1123		AU 1	999-	3820	8		1	9990	423	<
	7531				В2		2002	1010										
BR	9910	177			А		2001	0109		BR 1	999-	1017	7		1	9990	423	<
EP	1083	792			A1		2001	0321		EP 1	999-	9207	48		1	9990	423	<
EP	1083	792			В1		2003	0924										
	R:	ΑT,	BE,	CH,	DE,	DK,												
	2001						2001	1028			001-							
	2002						2002				000-							<
	5085				Α		2003				999-							
	2503				Τ		2003				999-							
	1083				_			0227			999-				_	9990		
	2934							0414			-000					9990		
	2204							0416			999-				_	9990		
	2846							0804			-000					9990		
	1392 5816	_			A B			0831			.999- .999-		_			9990 9990		
	2000		572								:999- :000-:							
	6436						2001				:000-:					0001		\
	APP	-			DI		2002	0020		DE 1	.000- .998- .999-:	1981	9628	1	A 1 W 1	9980	504	

OTHER SOURCE(S): MARPAT 131:318921

GI

The title mixts. comprise a carbamate I (T = CH or N; n = 0, 1 or 2; R = halo, C1-4 alkyl or alkyl halide), the oxime ether carboxylic acid ester II or the oxime ether carboxylic acid amide III and IV [R1 = (un)substituted C1-4 alkyl, C2-4 alkenyl, C2-4 alkynyl or alkylcycloalkyl; R2 = C1-4 alkyl or alkyl halide; R3 = H, halo, C1-4 alkyl, alkoxy, alkylthio, alkylamino, alkyl halide or haloalkoxy; Y = 0, S, CHR4 or NR5; R4, R5 = R2; n = 0, 1, 2 or 3].

II 175013-18-0 175013-22-6

RL: AGR (Agricultural use); BIOL (Biological study); USES (Uses) (mixture containing; synergistic fungicide)

RN 175013-18-0 ZCAPLUS

CN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

$$\begin{array}{c|c}
 & \text{N} & \text{N} & \text{O-CH}_2 \\
 & \text{MeO-C-N} \\
 & \text{MeO$$

RN 175013-22-6 ZCAPLUS

CN Carbamic acid, methoxy[2-[[[1-(4-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 41 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1999:640833 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 131:257438

TITLE: Preparation of 3-aralkylidene-2-oxopyrrole-3-

carboxylates as crop protection agents.

INVENTOR(S): Wagner, Oliver; Otten, Martina; Westphalen, Karl-otto;

Walter, Helmut; Harries, Volker

PATENT ASSIGNEE(S): Basf A.-G., Germany SOURCE: PCT Int. Appl., 51 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA.	TENT	NO.			KIN	D	DATE			APPL	ICAT	ION 1	.OV		D.	ATE		
WO	9950	 243			A1	_	1999	1007	,	WO 1	 999-:	EP20	 06		1	 9990	324	<
	W:	AL,	ΑU,	BG,	BR,	BY,	CA,	CN,	CZ,	GE,	HU,	ID,	IL,	IN,	JP,	KR,	KΖ,	
		LT,	LV,	MK,	MX,	NO,	NΖ,	PL,	RO,	RU,	SG,	SI,	SK,	TR,	UA,	US,	ZA,	
		ΑM,	AZ,	BY,	KG,	KΖ,	MD,	RU,	ТJ,	TM								
	RW:	ΑT,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	ΙΤ,	LU,	MC,	NL,	
		PT,	SE															
CA	2325	904			A1		1999	1007	1	CA 1	999-	2325	904		1	9990	324	<
AU	9937	020			Α		1999	1018		AU 1	999-	3702	0		1	9990	324	<
EP	1066	256			A1		2001	0110		EP 1	999-	9191	37		1	9990	324	<
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	NL,	SE,	PT,	FΙ		
JP	2002	5099	17		Τ		2002	0402	1	JP 2	000-	5411	48		1	9990	324	<
US	6548	451			В1		2003	0415		US 2	001-	6470	10		2	0010	124	
PRIORIT	Y APP	LN.	INFO	.:						DE 1	998-	1981	4040		A 1	9980	330	
									,	WO 1	999-:	EP20	06	1	W 1	9990	324	
OTHER SO	OURCE	(S):			MAR	PAT	131:	2574.	38									

GΙ

AB Use of title compds. [I; R1 = (substituted) (condensed) aryl, heteroaryl; R2 = alkyl, cycloalkyl, (substituted) aryl, heteroaryl; A = CO2R3, CONR3R4; R3, R4 = H, (substituted) alkyl, alkenyl, cycloalkyl, alkylaryl] as crop protection agents is claimed. Thus, Et 2-methyl-5-oxo-4,5- dihydropyrrole-3-carboxylate and 2-ethylbenzaldehyde were stirred with cat. HCl in EtOH to give Et E/Z-4-(2-ethylbenzylidene)-2-methyl-5-oxo-4,5- dihydropyrrole-3-carboxylate. Several I at 3 g/ha postemergent gave complete control of Sinapis alba and Setaria italica.

IT 244300-39-8P 244300-41-2P 244300-71-8P

244300-73-0P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic

preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)
 (preparation of 3-aralkylidene-2-oxopyrrole-3-carboxylates as crop
 protection agents)

RN 244300-39-8 ZCAPLUS

CN 1H-Pyrrole-3-carboxylic acid, 4-[[1-(4-chlorophenyl)-1H-pyrazol-4-yl]methylene]-4,5-dihydro-2-methyl-5-oxo-, methyl ester, (4Z)- (9CI) (CF INDEX NAME)

Double bond geometry as shown.

RN 244300-41-2 ZCAPLUS

CN 1H-Pyrrole-3-carboxylic acid, 4-[[3-(4-chlorophenyl)-5-isoxazolyl]methylene]-4,5-dihydro-2-methyl-5-oxo-, methyl ester, (4Z)-(9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 244300-71-8 ZCAPLUS

CN 1H-Pyrrole-3-carboxylic acid, 4-[[1-(4-chlorophenyl)-1H-pyrazol-4-yl]methylene]-4,5-dihydro-2-methyl-5-oxo-, ethyl ester, (4Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 244300-73-0 ZCAPLUS

CN 1H-Pyrrole-3-carboxylic acid, 4-[[3-(4-chlorophenyl)-5-isoxazolyl]methylene]-4,5-dihydro-2-methyl-5-oxo-, ethyl ester, (4Z)-(9CI) (CA INDEX NAME)

Double bond geometry as shown.

$$\begin{array}{c} \text{Cl} \\ \\ \\ \\ \\ \\ \\ \\ \end{array} \begin{array}{c} \text{OEt} \\ \\ \\ \\ \\ \\ \end{array}$$

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 42 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1999:620484 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 131:243076

TITLE: Preparation of hydroxyanilines as herbicides

INVENTOR(S): Sato, Kazuo; Sano, Hiroki; Komai, Hiroyuki; Kudou,

Noriaki; Morimoto, Soji; Kadotani, Junji

PATENT ASSIGNEE(S): Sankyo Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 43 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Fatent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 11263775	А	19990928		19980907 <
PRIORITY APPLN. INFO.:			JP 1997-242967 A	19970908
OTHER SOURCE(S):	MARPAT	131:243076		

$$\begin{array}{c} R^3 \\ \text{CHQ} \\ \\ R^1 \text{CONH} \end{array}$$

Title compds. I (R1 = alkoxy; R2 = alkyl, cycloalkyl, alkoxy, halo; R3 = H, alkyl; Q = heterocyclyl, except oxazolyl, 2-benzoxazolyl, thiazolyl, 2-benzothiazolyl) and their salts, useful as herbicides, are prepared Thus, reaction of 2-methyl-4-hydroxyaniline with 5-chloro-2- chloromethylthiophene in DMF in the presence of NaH gave 81.6% 4-(5-chlorothiophen-2-ylmethoxy)-2-methylaniline, reaction of which with Me chloroformate in CH2Cl2 in the presence of 4-dimethylaminopyridine gave 92.3% Me [4-(5-chlorothiophen-2-ylmethoxy)-2-methylphenyl]carbamate (II). II showed herbicidal activity at 20 g/are against Echinocloa crus-galli with no toxicity to rice.

IT 244175-43-7P 244175-44-8P 244175-45-9P 244175-46-0P 244175-52-8P 244175-57-3P 244175-58-4P 244175-59-5P 244175-61-9P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of hydroxyanilines as herbicides)

RN 244175-43-7 ZCAPLUS

CN Carbamic acid, [4-[(3-phenyl-5-isoxazolyl)methoxy]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 244175-44-8 ZCAPLUS

CN Carbamic acid, [2-methyl-4-[(3-phenyl-5-isoxazolyl)methoxy]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 244175-45-9 ZCAPLUS

CN Carbamic acid, [4-[[3-(2-chlorophenyl)-5-methyl-4-isoxazolyl]methoxy]-2-methylphenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 244175-46-0 ZCAPLUS

CN Carbamic acid, [4-[[3-(2,6-dichlorophenyl)-5-methyl-4-isoxazolyl]methoxy]-2-methylphenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 244175-52-8 ZCAPLUS

CN Carbamic acid, [2-methyl-4-[(5-methyl-1-phenyl-1H-pyrazol-3-yl)methoxy]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 244175-57-3 ZCAPLUS

CN Carbamic acid, [2-methyl-4-[(3-methyl-1-phenyl-1H-pyrazol-5-yl)methoxy]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 244175-58-4 ZCAPLUS

CN Carbamic acid, [2-methyl-4-[[1-phenyl-3-(trifluoromethyl)-1H-pyrazol-5-yl]methoxy]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

Ph
$$CH_2-O$$
 Me $NH-C-OMe$

RN 244175-59-5 ZCAPLUS

CN Carbamic acid, [2-methyl-4-[[3-(pentafluoroethyl)-1-phenyl-1H-pyrazol-5-yl]methoxy]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

Ph
$$CH_2 - O$$
 $NH - C - OMe$

RN 244175-61-9 ZCAPLUS

CN Carbamic acid, [4-[[1-(4-chlorophenyl)-5-(trifluoromethyl)-1H-pyrazol-4-yl]methoxy]-2-methylphenyl]-, methyl ester (9CI) (CA INDEX NAME)

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PAGE 2-A

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L89 ANSWER 43 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1999:312718 ZCAPLUS Full-text

DOCUMENT NUMBER: 131:5260

TITLE: Preparation of azole ring-containing phenylcarboxylic

acids as lipid formation inhibitors

INVENTOR(S): Kitaide, Makoto; Ono, Tomoyasu; Terada, Tadashi; Asao,

Tetsuji; Yamamoto, Akiyoshi; Yamada, Haruo; Miyake,

Hidekazu

PATENT ASSIGNEE(S): Taiho Pharmaceutical Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 29 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 11130753	A	19990518	JP 1997-300384	19971031 <
JP 3694774	B2	20050914		
PRIORITY APPLN. INFO.:			JP 1997-300384	19971031
OTHER SOURCE(S):	MARPAT	131:5260		
GI				

The title compds. I [A = lower alkyl, (un) substituted Ph, (un) substituted pyridyl, in which the substituent is halo, lower alkyl, lower alkoxy, alkylamino; Q = imidazolyl, triazolyl, pyrazolyl, thiazolyl which may be substituted with lower alkyl, etc.; B = 0, NR7 (R7 = H, lower alkyl); R1 = H, halo, lower alkoxy; R2 = H, lower alkyl; n = 1, 2; m = 0, 1] or their salts are prepared I or their salts inhibit fatty acid synthesis and cholesterol synthesis and are useful as hypolipemics. A THF solution of 1-(4-chlorophenyl)-5-methyl-4-hydroxymethylpyrazole (preparation given) was treated with SOC12 and the resulting 1-(4-chlorophenyl)-5-methyl-4-chloromethylpyrazole was treated with p-HOC6H4CO2Me to give 1-(4-chlorophenyl)-5-methyl-4-(4'-methoxycarbonylphenoxy)methylpyrazole. Similarly prepared 1-phenyl-5-methyl-4-(4'-methoxycarbonylphenoxy)methylpyra zole significantly lowered serum triglycerides and VLDL cholesterol.

IT 225930-55-2P 225930-56-3F 225930-59-6P 225930-67-6P 225930-68-7P 225930-69-8P 225930-70-1F 225930-72-3P 225930-73-4P 225930-77-8P 225930-75-6P 225930-80-3P 225930-81-4P 225930-82-5P 225930-83-6P 225930-84-7P 225930-86-9P 225930-87-0P 225930-88-1P 225930-90-5F 225930-91-6P

225930-92-7P 225930-94-9P 225930-95-0P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of azole ring-containing phenylcarboxylic acids as lipid formation $\ensuremath{\mathsf{S}}$

inhibitors)

RN 225930-55-2 ZCAPLUS

CN Benzoic acid, 4-[[1-(4-chlorophenyl)-5-methyl-1H-pyrazol-4-yl]methoxy]-, methyl ester (9CI) (CA INDEX NAME)

RN 225930-56-3 ZCAPLUS

CN Benzoic acid, 4-[[[1-(4-chlorophenyl)-5-methyl-1H-pyrazol-4-

yl]methyl]methylamino]-, ethyl ester (9CI) (CA INDEX NAME)

PAGE 1-A

$$\begin{array}{c} \text{C1} \\ \text{N} \\ \text{N} \\ \text{Me} \\ \\ \text{C} \\ \text{OEt} \\ \end{array}$$

PAGE 2-A

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RN 225930-59-6 ZCAPLUS

CN Benzoic acid, 4-[[5-methyl-1-(4-pyridinyl)-1H-pyrazol-4-yl]methoxy]-, methyl ester (9CI) (CA INDEX NAME)

RN

CN Benzoic acid, 4-[[1-(4-chlorophenyl)-5-methyl-1H-pyrazol-4-yl]methoxy]- (9CI) (CA INDEX NAME)

RN 225930-68-7 ZCAPLUS

CN Benzoic acid, 4-[(5-methyl-1-phenyl-1H-pyrazol-4-yl)methoxy]-, methyl ester (9CI) (CA INDEX NAME)

RN 225930-69-8 ZCAPLUS

CN Benzoic acid, 4-[[1-(4-chlorophenyl)-1H-pyrazol-4-yl]methoxy]-, methyl ester (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

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RN 225930-70-1 ZCAPLUS

CN Benzoic acid, 4-[[1-(4-chlorophenyl)-5-methyl-1H-pyrazol-4-yl]methoxy]-, ethyl ester (9CI) (CA INDEX NAME)

RN 225930-72-3 ZCAPLUS

CN Benzoic acid, 4-[[5-methyl-1-(2-pyridinyl)-1H-pyrazol-4-yl]methoxy]-, methyl ester (9CI) (CA INDEX NAME)

RN 225930-73-4 ZCAPLUS

CN Benzoic acid, 4-[[1-(4-methoxyphenyl)-5-methyl-1H-pyrazol-4-yl]methoxy]-, methyl ester (9CI) (CA INDEX NAME)

RN 225930-74-5 ZCAPLUS

CN Benzoic acid, 4-[[3-(dimethylamino)-1-phenyl-1H-pyrazol-4-yl]methoxy]-, methyl ester (9CI) (CA INDEX NAME)

RN 225930-75-6 ZCAPLUS

CN Benzoic acid, 4-[[1-(4-chlorophenyl)-5-(dimethylamino)-1H-pyrazol-4-yl]methoxy]-, methyl ester (9CI) (CA INDEX NAME)

RN 225930-76-7 ZCAPLUS

CN Benzoic acid, 4-[[1-(4-chlorophenyl)-5-methyl-1H-pyrazol-4-yl]methoxy]-3-methoxy-, methyl ester (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

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RN 225930-77-8 ZCAPLUS

CN Benzoic acid, 3-chloro-4-[[1-(4-chlorophenyl)-5-methyl-1H-pyrazol-4-yl]methoxy]-, ethyl ester (9CI) (CA INDEX NAME)

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PAGE 2-A

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RN 225930-78-9 ZCAPLUS

CN Benzoic acid, 3-chloro-4-[(5-methyl-1-phenyl-1H-pyrazol-4-yl)methoxy]-, ethyl ester (9CI) (CA INDEX NAME)

RN 225930-80-3 ZCAPLUS

CN Benzoic acid, 4-[[1-(4-chlorophenyl)-1H-pyrazol-4-yl]methoxy]- (9CI) (CA INDEX NAME)

RN 225930-81-4 ZCAPLUS

CN Benzoic acid, 4-[[1-[4-(dimethylamino)phenyl]-5-methyl-1H-pyrazol-4-yl]methoxy]-, ethyl ester (9CI) (CA INDEX NAME)

RN 225930-82-5 ZCAPLUS

CN Benzoic acid, 4-[[1-methyl-5-(1H-pyrrol-1-yl)-1H-pyrazol-4-yl]methoxy]-, methyl ester (9CI) (CA INDEX NAME)

RN 225930-83-6 ZCAPLUS

CN Benzoic acid, 4-[[5-methyl-1-(4-methylphenyl)-1H-pyrazol-4-yl]methoxy]-, ethyl ester (9CI) (CA INDEX NAME)

RN 225930-84-7 ZCAPLUS

CN Benzoic acid, 4-[[1-(4-fluorophenyl)-5-methyl-1H-pyrazol-4-yl]methoxy]-, ethyl ester (9CI) (CA INDEX NAME)

RN 225930-86-9 ZCAPLUS

CN Benzoic acid, 4-[[5-methyl-1-(2-pyridinyl)-1H-pyrazol-4-yl]methoxy]- (9CI) (CA INDEX NAME)

RN 225930-87-0 ZCAPLUS

CN Benzoic acid, 4-[[1-(4-methoxyphenyl)-5-methyl-1H-pyrazol-4-yl]methoxy]- (9CI) (CA INDEX NAME)

RN 225930-88-1 ZCAPLUS
CN Benzoic acid, 4-[[5-methyl-1-(4-pyridinyl)-1H-pyrazol-4-yl]methoxy]- (9CI)
(CA INDEX NAME)

RN 225930-90-5 ZCAPLUS
CN Benzoic acid, 4-[[1-(4-chlorophenyl)-5-methyl-1H-pyrazol-4-yl]methoxy]-3-methoxy- (9CI) (CA INDEX NAME)

RN 225930-91-6 ZCAPLUS

CN Benzoic acid, 3-chloro-4-[[1-(4-chlorophenyl)-5-methyl-1H-pyrazol-4-yl]methoxy]- (9CI) (CA INDEX NAME)

RN 225930-92-7 ZCAPLUS

CN Benzoic acid, 3-chloro-4-[(5-methyl-1-phenyl-1H-pyrazol-4-yl)methoxy]- (9CI) (CA INDEX NAME)

RN 225930-94-9 ZCAPLUS

CN Benzoic acid, 4-[[[1-(4-chlorophenyl)-5-methyl-1H-pyrazol-4-yl]methyl]methylamino]- (9CI) (CA INDEX NAME)

RN 225930-95-0 ZCAPLUS

CN Benzoic acid, 4-[[[1-(4-chlorophenyl)-5-methyl-1H-pyrazol-4-yl]methoxy]methyl]- (9CI) (CA INDEX NAME)

L89 ANSWER 44 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1999:181680 ZCAPLUS Full-text

DOCUMENT NUMBER: 130:209706

TITLE: Preparation of N-acylated

pyrazolyloxymethylphenylhydroxylamines and related

compounds.

INVENTOR(S): Klintz, Ralf; Goetz, Norbert; Keil, Michael; Heilig,

Manfred; Wingert, Horst; Vogelbacher, Uwe Josef; Wahl,

Josef; Witterich, Frank

PATENT ASSIGNEE(S): BASF A.-G., Germany SOURCE: Ger. Offen., 10 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Fatest LANGUAGE: German

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PA'	IENT 1	NO.			KINI	D I	DATE			APPL	ICAT:	I NOI	.OV		D.	ATE		
DE	1973	8864			A1	-	1999	0311		DE 1	997-1	1973	8864		1	9970	905	<
CA	2302	937			A1	_	1999	0318	(CA 1	998-2	23029	937		1	9980	821	<
WO	9912	911			A1	_	1999	0318	1	WO 1	998-1	EP533	32		1	9980	821	<
	W:	AL,	ΑU,	BG,	BR,	BY,	CA,	CN,	CZ,	GE,	HU,	ID,	IL,	JP,	KR,	KΖ,	LT,	
		LV,	MK,	MX,	NO,	NZ,	PL,	RO,	RU,	SG,	SI,	SK,	TR,	UA,	US,	AM,	ΑZ,	
		KG,	MD,	ΤJ,	TM													
	RW:	ΑT,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	ΙT,	LU,	MC,	NL,	
		PT,	SE															
AU	9892	643			A	-	1999	0329		AU 1	998-9	92643	3		1	9980	821	<
EP	1012	144			A1	2	2000	0628		EP 1	998-9	9452	76		1	9980	821	<
EP	1012	144			В1	2	2003	0312										
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	NL,	SE,	PT,	IE,	FΙ	
BR	9812	041			Α	2	2000	0926	:	BR 1	998-1	1204	1		1	9980	821	<
HU	2000	0406	3		A2	2	2001	0328		HU 2	000-	4063			1	9980	821	<
JP	2001	5158	90		Τ	2	2001	0925		JP 2	000-	51072	20		1	9980	821	<
AT	2342	89			Τ	2	2003	0315		AT 1	998-9	9452	76		1	9980	821	
CN	1117	080			В	2	2003	0806	(CN 1	998-8	8094	44		1	9980	821	
ES	2195	388			Т3	2	2003	1201			998-9					9980	821	
CZ	2970	14			В6	2	2006	0816	(CZ 2	000-	764			1	9980	821	
US	6255	489			В1	2	2001	0703	1	US 2	000-	48650	00		2	0000	229	<
PRIORIT	Y APP	LN.	INFO	.:						DE 1	997-1	1973	8862		A 1	9970	905	
										DE 1	997-1	1973	8864		A 1	9970	905	

WO 1998-EP5332 W 19980821

OTHER SOURCE(S): CASREACT 130:209706; MARPAT 130:209706

GΙ

AΒ Title compds. (I; R1 = alkoxycarbonyl, alkylcarbonyl, alkylaminocarbonyl, dialkylaminocarbonyl; R2 = H, alkyl; R3 = H, halo, cyano, alkyl, haloalkyl, alkoxy, alkoxycarbonyl, AB, etc.; R4 = halo, alkyl, haloalkyl, alkoxycarbonyl; X = N, CH; A = O, CH2, OCH2, CH2O2C, CH:CH, CH:NO, etc.; B = Ph, naphthyl, pyridinyl, pyrazinyl, pyrimidinyl, pyrazolyl, imidazolyl, oxazolyl, isoxazolyl, etc.; n = 0-3), were prepared by hydrogenation of the corresponding nitro compds. in a mixture of an aprotic solvent and an aliphatic amine followed by N-acylation of the resulting unisolated hydroxylamine and optional O-alkylation. Thus, 2-[N-(pchlorophenyl)pyrazolyl-3- oxymethyl]nitrobenzene was hydrogenated over Pt/C in PhMe/PrNH2 at 5° and 100 bar H2 for 2 h; PrNH2 was distilled off to give 93.4% N-hydroxy-N-2-[N-(p-chlorophenyl)pyrazolyl-3-oxymethyl]aniline as a PhMe solution C1CO2Me was added to a rapidly stirred emulsion of the above solution and H2O over 2 h followed by 2.5 h stirring at 35° to give 88% Nhydroxy-N-[2-[N-(p-chlorophenyl)pyrazolyl-3-oxymethyl]phenyl]carbamic acid Me ester.

IT 220897-48-3P 220897-58-5P 220897-76-7P 220897-80-3P 220897-86-9P 220897-91-6P 220897-96-1P 220898-10-2P 220898-33-9P

RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)

(preparation of N-acylated azolyloxymethylphenylhydroxylamines and related compds.)

RN 220897-48-3 ZCAPLUS

CN Carbamic acid, hydroxy[2-[[(1-phenyl-1H-pyrazol-3-yl)oxy]methyl]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{N} & \text{OOH} \\ \text{MeO-C-N} \\ \text{O-CH}_2 \end{array}$$

RN 220897-58-5 ZCAPLUS

CN Urea, N-hydroxy-N'-methyl-N-[2-[[(1-phenyl-1H-pyrazol-3-yl)oxy]methyl]phenyl]- (9CI) (CA INDEX NAME)

RN 220897-76-7 ZCAPLUS

CN Carbamic acid, [2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]hydroxy-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{C1} & \begin{array}{c} \text{OOH} \\ \text{MeO-C-N} \end{array} \end{array}$$

RN 220897-80-3 ZCAPLUS

CN Urea, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-hydroxy-N'-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \circ & \circ H \\ \text{MeNH-C-N} \\ & \circ - \circ H_2 \end{array}$$

RN 220897-86-9 ZCAPLUS

CN Carbamic acid, [2-[[[1-(4-chlorophenyl)-5-methyl-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 220897-91-6 ZCAPLUS

CN Carbamic acid, [2-[[[1-cyclohexyl-4-(trifluoromethyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 220897-96-1 ZCAPLUS

CN Carbamic acid, [2-[[[4-chloro-1-(5-chloro-2-pyridinyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} C1 & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

RN 220898-10-2 ZCAPLUS

CN Urea, N-methoxy-N'-methyl-N-[2-[[(1-phenyl-1H-pyrazol-3-yl)oxy]methyl]phenyl]- (9CI) (CA INDEX NAME)

RN 220898-33-9 ZCAPLUS

CN Urea, N-[2-[[[1-(2,4-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-N-methoxy-N'-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & &$$

L89 ANSWER 45 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1999:181678 ZCAPLUS Full-text

DOCUMENT NUMBER: 130:222827

TITLE: Preparation of N-aryl- and N-heterocyclyl-

hydroxylamines

INVENTOR(S): Klintz, Ralf; Heilig, Manfred; Keil, Michael;

Vogelbacher, Uwe Josef; Wahl, Josef; Wingert, Horst;

Goetz, Norbert; Daun, Gregor

PATENT ASSIGNEE(S): BASF A.-G., Germany SOURCE: Ger. Offen., 14 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

GΙ

PA:	TENT NO.		KIN	D DA'	ΓE	Al	PPLI	CATI	ION :	NO.		D	ATE		
CA	19738862 2302937 9912911 W: AL, A		A1 A1	199 199	990318 990318	C2 W(A 19 O 19	98-2 98-E	2302 EP53	937 32		1 1	9980 9980	821 821	<
	LV, M KG, M			NZ, P	L, RO,	RU,	SG,	SI,	SK,	TR,	UA,	US,	AM,	AZ,	
	RW: AT, B: PT, S:	, ,	CY,	DE, Di	K, ES,	FI, I	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	
AU	9892643		Α	199	990329	Α	U 19	98-9	9264	3		1	9980	821	<
EP	1012144		A1	20	000628	El	P 19	98-9	9452	76		1	9980	821	<
EP	1012144		В1	20	030312										
	R: AT, B	E, CH,	DE,	DK, E	S, FR,	GB, (GR,	ΙΤ,	LI,	NL,	SE,	PT,	ΙE,	FΙ	
BR	9812041		Α	20	000926	BI	R 19	98-1	1204	1		1	9980	821	<
	200004063				010328							1	9980	821	<
JP	2001515890		T	20	010925	J1	P 20	00 - 5	5107	20		1	9980	821	<
AT	234289		T	20	030315	A.	T 19	98-9	9452	76		1	9980	821	
CN	1117080		В	20	030806	CI	N 19	98-8	3094	44		1	9980	821	
ES	2195388		Т3	20	031201	E	S 19	98-9	9452	76		1	9980	821	
CZ	297014		В6	20	060816	C:	Z 20	00-	764			1	9980	821	
US	6255489		В1	20	010703	U:	S 20	00 - 4	4865	00		2	0000	229	<
MX	200002189		А	20	001020	M	X 20	00 - 2	2189			2	0000	302	<
PRIORIT	Y APPLN. IN	· · · ·				DI	E 19	97-1	1973	8862		A 1	9970	905	
						DI	E 19	97-1	1973	8864		A 1	9970	905	
										32		W 1	9980	821	
OTHER SO	OURCE(S):		CAS:	REACT :	130 : 22	2827;	MAR	PAT	130	:222	827				

AB Aromatic and heteroarom. nitro compds. are reduced to the hydroxylamines by treatment with an amine in presence of a transition metal catalyst in inert aprotic solvent. Thus, the nitro compound I [R = NO2] was treated with H in PhMe in presence of Pt-C and PrNH2 to give 93.4% I [R = NHOH] as a solution in PhMe which was treated with ClCO2Me in aqueous PhMe to give I [R = N(OH)CO2Me] in 88% overall yield.

IT 220897-76-7F

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of N-aryl- and N-heterocyclyl-hydroxylamines)

RN 220897-76-7 ZCAPLUS

CN Carbamic acid, [2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]hydroxy-, methyl ester (9CI) (CA INDEX NAME)

L89 ANSWER 46 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1999:7973 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 130:52416

TITLE: Pesticidal 1-aryl-3-iminopyrazoles INVENTOR(S): Manning, David Treadway; Wu, Tai-teh

PATENT ASSIGNEE(S): Rhone-Poulenc Agro, Fr. SOURCE: PCT Int. Appl., 70 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	TENT																	
	9856																	<
	W:	AL,	ΑU,	BA,	BB,	BG,	BR,	CA,	CN,	CU,	CZ,	EE,	GE,	GW,	HU,	ID,	IL,	,
		IS,	JP,	KP,	KR,	LK,	LR,	LT,	LV,	MG,	MK,	MN,	MX,	NO,	NΖ,	PL,	RO,	,
		SG,	SI,	SK,	SL,	TR,	TT,	UA,	UZ,	VN,	YU,	AM,	ΑZ,	BY,	KG,	KΖ,	MD,	,
		RU,	ТJ,	TM														
	RW:	GH,	GM,	ΚE,	LS,	MW,	SD,	SZ,	UG,	ZW,	ΑT,	BE,	CH,	DE,	DK,	ES,	FI,	,
		FR,	GB,	GR,	IE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,	,
							SN,											
ZA	9801	934			Α		1999	0906		ZA 1	998-	1934			1	9980	306	<
CA	2283	465			A1		1998:	1217		CA 1	998-	2283	465		1	9980	309	<
ΑU	9870	415			Α		1998:	1230		AU 1	998-	7041	5		1	9980	309	<
	7450																	
	5965																	
BR	9808	019														9980	309	<
EE	9900	402			А		2000	0417		EE 1	999-	402			1	9980	309	<
	4014																	
EP	1007	513			A1		2000	0614		EP 1	998-	9170	82		1	9980	309	<
	R:						FR,	GB,	GR,	ΙT,	LI,	LU,	NL,	SE,	MC,	PT,	IE,	
					FI,													
TR	9902	211			Т2		2000	0621				2211				9980		
HU	2000	0192	3		A2		2001	0129								9980		
	2001									JP 1	998-	5463	87		1	9980	309	<
	3785																	
CN	1107	673			В		2003	0507		CN 1	998-	8039	53		1	9980	309	
ΑP	1158				Α		2003	0630		AP 1	999-	1645			1	9980	309	
	W:																	
CZ	2961	62			В6		2006	0111		CZ 1	999-	3184			1	9980	309	

TW	486470	В	20020511	TW	1998-87103503		19980310	<
NO	9904355	A	19991110	NO	1999-4355		19990908	<
NO	313828	B1	20021209					
MX	9908352	A	20000228	MX	1999-8352		19990910	<
BG	103775	A	20010531	ВG	1999-103775		19991004	<
BG	64128	B1	20040130					
HK	1025320	A1	20040116	HK	2000-104482		20000720	
PRIORITY	APPLN. INFO.:			US	1997-40135P	Р	19970310	
				WO	1998-EP1764	W	19980309	
OTHER SC	OURCE(S):	MARPAT	130:52416					

GΙ

AΒ The title compds. [I; R31 = H, CN, NO2, etc.; R32 = C1-6 alkyl, C3-7 cycloalkyl, etc.; R33 = a lone pair of electrons, O, S, etc.; R4 = C1-6 alkyl, C3-6 cycloalkyl, C4-8 (cycloalkyl)alkyl, etc.; R5 = H, halo, CN, etc.; Z = N, CH, C(halo), etc.; R12-R15 = H, halo, CN, etc.], useful as pesticides, especially for controlling arthropods, or as intermediates to other pesticides, were prepared Thus, reaction of 3-acetyl-5-amino-1-[2,6dichloro-4-(trifluoromethyl)phenyl]-4-methylsulfinyl-1H-pyrazole with aniline in the presence of p-TsOH in C6H6 afforded I [R32 = Me; R31 = Ph; R33 = a lone pair of electrons; R4 = MeS(O); R5 = NH2; R12 = C1, R13 = R15 = H; R14 = CF3; Z = C(Cl)] which showed high systemic activity on aphids and on greenbugs. ΙT 217437-17-72

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (pesticidal 1-aryl-3-iminopyrazoles)

217437-17-7 ZCAPLUS RN

Benzamide, 3-[[1-[5-amino-1-[2,6-dichloro-4-(trifluoromethyl)phenyl]-4-CN (methylsulfinyl)-1H-pyrazol-3-yl]ethylidene]amino]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} F_3C & & \\ \hline & N & \\ \hline & C-NH_2 \\ \hline & \\ & & \\ \hline \end{array}$$

REFERENCE COUNT: THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 47 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1998:793053 ZCAPLUS Full-text

DOCUMENT NUMBER: 130:34479

TITLE: Synergistic fungicidal mixtures

INVENTOR(S): Schelberger, Klaus; Scherer, Maria; Saur, Reinhold;

Sauter, Hubert; Muller, Bernd; Birner, Erich; Leyendecker, Joachim; Ammermann, Eberhard; Lorenz,

Gisela; Strathmann, Siegfried

PATENT ASSIGNEE(S): Basf Aktiengesellschaft, Germany; et al.

SOURCE: PCT Int. Appl., 20 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Fatent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

P	PATEN	IT I	. O <i>l</i> .			KIN	D	DATE			APPL	ICAT	ION :	NO.		D.	ATE		
W	70 98	53	 693			A1	_	 1998	1203	,	WO 1	 998-:	EP29	 13		1	9980	 518 <	:
	M	:	AL,	ΑU,	BG,	BR,	BY,	CA,	CN,	CZ,	GE,	HU,	ID,	IL,	JP,	KR,	KΖ,	LT,	
			LV,	MX,	NO,	NZ,	PL,	RO,	RU,	SG,	SI,	SK,	TR,	UA,	US,	AM,	ΑZ,	KG,	
			MD,	ТJ,	TM														
	R	: W	ΑT,	BE,	CH,	CY,	DE,	DK,	ES,	FΙ,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	
			PT,	SE															
A	4U 98	79:	132			Α		1998	1230		AU 1	998-	7913	2		1	9980	518 <	:
I	N 19	981	MA01	166		Α		2005	0304		IN 1	998-	MA11	66		1	9980	529	
PRIORI	TY A	PP:	LN.	INFO	.:						DE 1	997-	1972	2652		A 1	9970	530	
										,	WO 1	998-	EP29	13	1	W 1	9980	518	
OTHER	COLLD		101			MAD		120.	2447	n									

OTHER SOURCE(S): MARPAT 130:34479

GΙ

- AB The title mixts. comprise a carbamate I (X = CH or N; n = 0, 1 or 2; R = H, halo, C1-4 alkyl or haloalkyl) or the oxime ether carboxamide II and a fungicidal copper (II) compound
- IT 216659-76-6

RL: AGR (Agricultural use); BIOL (Biological study); USES (Uses) (synergistic fungicide)

RN 216659-76-6 ZCAPLUS

CN Carbamic acid, [2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester, mixt. with copper hydroxide sulfate (Cu4(OH)6(SO4)) (9CI) (CA INDEX NAME)

CM 1

CRN 175013-18-0 CMF C19 H18 C1 N3 O4

CM 2

CRN 1333-22-8

CMF Cu . H O . O4 S

CCI TIS

CM 3

CRN 14808-79-8

CMF 04 S

CM 4

CRN 14280-30-9

CMF H O

OH-

CM 5

CRN 7440-50-8

CMF Cu

Cu

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 48 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1998:793051 ZCAPLUS Full-text

DOCUMENT NUMBER: 130:34477

TITLE: Synergistic fungicidal mixtures

INVENTOR(S): Schelberger, Klaus; Scherer, Maria; Saur, Reinhold;

Sauter, Hubert; Muller, Bernd; Birner, Erich; Leyendecker, Joachim; Ammermann, Eberhard; Lorenz,

Gisela; Strathmann, Siegfried

PATENT ASSIGNEE(S): Basf Aktiengesellschaft, Germany; et al.

SOURCE: PCT Int. Appl., 22 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	ENT						DATE			APPL	ICAT	ION	NO.		D.	ATE		
	9853						1998	1203		 WO 1	998-	 EP28	77		1	9980	515	<
	W:	AL,	ΑU,	BG,	BR,	BY,	CA,	CN,	CZ,	GE,	HU,	ID,	IL,	JP,	KR,	KΖ,	LT,	
		LV,	MX,	NO,	NZ,	PL,	RO,	RU,	SG,	SI,	SK,	TR,	UA,	US,	AM,	ΑZ,	KG,	
		MD,	ΤJ,	TM														
	RW:	ΑT,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	ΙT,	LU,	MC,	NL,	
		PT,	SE															
DE	1972	2225			A1		1998	1203		DE 1	997-	1972	2225		1	9970	528	<
CA	2289	786			A1		1998	1203		CA 1	998-	2289	786		1	9980	515	<
ΑU	9880	178			Α		1998	1230		AU 1	998-	8017	8		1	9980	515	<
EP	9846	95			A1		2000	0315		EP 1	998-	9282	74		1	9980	515	<
EP	9846	95			В1		2002	0327										
	R:									GR,	ΙT,	LI,	NL,	SE,	PT,	IE,	SI,	FΙ
	9809	480			Α		2000	0620		BR 1	998-	9480			1	9980	515	<
HU	2000 2001	0207	7		A2		2000	1028			000-							
JΡ	2001	5267	00		Τ		2001	1218			999-							
	5009						2002	0201		NZ 1	998-	5009	44		1	9980	515	<
	2148						2002	0415			998-							
TW	4101	42			В		2000	1101		TW 1	998-	8710	7672		1	9980	518	<
ΙN	1998	MA01	120		Α		2005	0304			998-					9980		
ZA	9804	508			Α		1999	1129			998-							
MX	9910	160			Α		2000	0430		MX 1	999-	1016	0		1	9991	105	<
US	6258	801			В1		2001	0710		US 1	999-	4234	62		1	9991	109	<
RITY	APP	LN.	INFO	.:						DE 1	997-	1972	2225		A 1	9970	528	
										WO 1	998-	EP28	77	,	W 1	9980	515	
D 00	TIDAD	(0)				m	100	0 4 4 5	_									

OTHER SOURCE(S): MARPAT 130:34477

GΙ

$$\underset{MeO_CO}{ } \overset{O}{\underset{OMe}{ }} \overset{T}{\underset{N}{ }} \overset{}{\underset{N}{ }} \overset{R_{n}}{\underset{I}{ }}$$

The title mixts. contain a carbamate I (T = CH or N; n = 0, 1 or 2; R = halo, C1-4 alkyl or halo alkyl) and a phosphonate R2OPH(O)OY [Y = H, group I, II or III metal or NR3R4R5R6; R2 = H, C1-18 alkyl, haloalkyl, nitroalkyl, (un)substituted C2-8 alkenyl or alkynyl, alkoxyalkyl, alkenylalkyl, (un)substituted aryl, cycloalkyl, alkylaryl or heterocyclyl with 5 or 6 ring atoms and N, O or S heteroatoms, whereby the heterocyclic group is linked to the O directly or via an aliphatic chain; R3-R6 = C1-4-alkyl or hydroxyalkyl].

II 216655-68-4

RL: AGR (Agricultural use); BIOL (Biological study); USES (Uses) (synergistic fungicide)

RN 216655-68-4 ZCAPLUS

CN Carbamic acid, [2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester, mixt. with aluminum tris(ethyl phosphonate) (9CI) (CA INDEX NAME)

CM 1

CRN 175013-18-0 CMF C19 H18 C1 N3 O4

CM 2

CRN 39148-24-8 CMF C2 H7 O3 P . 1/3 Al

0 HO_PH_OF+

●1/3 Al

L89 ANSWER 49 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1997:740080 ZCAPLUS Full-text

DOCUMENT NUMBER: 128:11111

TITLE: Synergistic fungicide mixtures

INVENTOR(S): Mueller, Bernd; Sauter, Hubert; Ammermann, Eberhard;

Lorenz, Gisela; Strathmann, Siegfried; Scherer, Maria;

Schelberger, Klaus; Leyendecker, Joachim

PATENT ASSIGNEE(S): BASF Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 20 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Fatent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

Р.	ATENT	NO.			KIN	D DATE	APPLICATION NO.		DATE
	974(W:	0686 AU,	BG,	BR,	A1 CA,	19971106 CN, CZ, GE,	WO 1997-EP2041 HU, IL, JP, KR, LV, AM, AZ, BY, KG, KZ,	MX,	19970423 < NO, NZ, PL,
	RW:	AT,	BE,	CH,	DE,	DK, ES, FI,	FR, GB, GR, IE, IT,	LU,	MC, NL, PT, SE
C.	A 2252	2511			A1	19971106	CA 1997-2252511		19970423 <
		7678			A	19971119	AU 1997-27678		19970423 <
A	U 7322	285			В2	20010412			
E	P 9000)19			A1	19990310	EP 1997-921700		19970423 <
E	P 9000)19			В1	20010829			
	R:	ΑT,	BE,	CH,	DE,	DK, ES, FR,	GB, GR, IT, LI, NL,	SE,	PT, IE, SI, FI
C	N 1216	5897			Α	19990519	CN 1997-194127		19970423 <
В	R 9708	3807			Α	19990803	BR 1997-8807 NZ 1997-332210		19970423 <
N	Z 3322	210			Α	20000228	NZ 1997-332210		19970423 <
J:	P 2000	05090	57		T	20000718	JP 1997-538544		
J.	P 3821	1486			В2	20060913			
A	T 204	707			Τ	20010915	AT 1997-921700		19970423 <
E	S 2163	3761			Т3	20020201			
P	T 9000)19			Τ	20020228			19970423 <
I.	N 199'	7MA00	834		Α	20050304	IN 1997-MA834		19970423
T.	W 4112	253			В	20001111			19970424 <
Z.	A 9703	3618			А	19990412	ZA 1997-3618		19970425 <
U	S 6172	2094							
	S 6239				В1	20010529			
G:	R 3036	5603			Т3	20011231			
PRIORI	TY API	PLN.	INFO	.:			DE 1996-19616716		A 19960426
							DE 1996-19617231		A 19960430
							DE 1996-19617234		A 19960430
							WO 1997-EP2041		
							US 1998-171649		A3 19981022

OTHER SOURCE(S): MARPAT 128:11111

GΙ

AB This invention concerns fungicide mixts. containing in a synergistically effective amount of a carbamate I [X = CH or N; n = 0, 1 or 2; R = halo or (halo) alkyl] and a dithiocarbamate selected from maneb, mancozeb, metiram and zineb, and/or cymoxanil.

IT 198956-59-1 198956-60-4 198956-62-6 198956-64-8

RL: AGR (Agricultural use); BIOL (Biological study); USES (Uses) (synergistic fungicide)

RN 198956-59-1 ZCAPLUS

CN Carbamic acid, [2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester, mixt. with 2-cyano-N-[(ethylamino)carbonyl]-2-(methoxyimino)acetamide (9CI) (CA INDEX NAME)

CM 1

CRN 175013-18-0 CMF C19 H18 C1 N3 O4

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ \end{array}$$

CM 2

CRN 57966-95-7 CMF C7 H10 N4 O3

RN 198956-60-4 ZCAPLUS

CN Carbamic acid, methoxy[2-[[[1-(4-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-, methyl ester, mixt. with 2-cyano-N-[(ethylamino)carbonyl]-2-(methoxyimino)acetamide (9CI) (CA INDEX NAME)

CM 1

CRN 175013-22-6 CMF C20 H21 N3 O4

CM 2

CRN 57966-95-7 CMF C7 H10 N4 O3

$$\begin{array}{c|c} & & & \circ & \text{N-OMe} \\ & & & & & \\ \text{EtNH-C-NH-C-C-CN} \end{array}$$

RN 198956-62-6 ZCAPLUS

CN Carbamic acid, [2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester, mixt. with metiram (9CI) (CA INDEX NAME)

CM 1

CRN 175013-18-0 CMF C19 H18 C1 N3 O4

CM 2

CRN 9006-42-2 CMF Unspecified CCI PMS, MAN

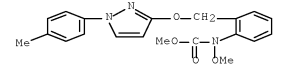
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

RN 198956-64-8 ZCAPLUS

CN Carbamic acid, methoxy[2-[[[1-(4-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-, methyl ester, mixt. with metiram (9CI) (CA INDEX NAME)

CM 1

CRN 175013-22-6 CMF C20 H21 N3 O4



CM 2

CRN 9006-42-2 CMF Unspecified CCI PMS, MAN

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

L89 ANSWER 50 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1997:303408 ZCAPLUS Full-text

DOCUMENT NUMBER: 126:273648

TITLE: Synergistic agrochemical fungicide comprising a

combination of an agent inhibiting respiration in the

cytochrome complex III and fenazaquin

INVENTOR(S): Bayer, Herbert; Sauter, Hubert; Ammermann, Eberhard;

Lorenz, Gisela; Strathmann, Siegfried; Koehle, Harald;

Retzlaff, Guenter

PATENT ASSIGNEE(S): BASF A.-G., Germany SOURCE: PCT Int. Appl., 57 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Fatent
LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	TENT NO.					DATE		AI	PPI	LICAT	ION :	NO.		DZ	ATE		
WO	9711606			A1	-												
	W: AU,																
	RW: AT,																
	401275																
IN	1996MA015					20050	0304	11	1]	1996-	MA15	29		19	9960!	902	
CA	2230888			A1	-	1997(0403	CZ	1	L996-	2230	888		19	9960	912	<
ΑU	9671288			A	-	19970	1417	ΑŪ	J 1	1996-	7128	8		19	9960	912	<
ΑU	711050			В2		19991	1007										
EP	862366			A1		19980	909	EI	?]	1996-	9325	15		19	9960	912	<
EP	862366			В1	2	20013	1121										
	R: AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB, (GR.	IT,	LI,	NL.	SE,	PT,	ΙE		
CN		,						CI								912	<
_								H									
_						20000			_								
	9610700			A		1999(RI	2 1	1996-	1070	Λ		1 (9960	912	<
	11511476			T		19991				1996-					9960		
	2158083			C2		20001				1998-					9960:		
	123631																
				A		20001				1996-		-			9960	-	
						20011				1996-					9960		
	9607963			A		19980				1996-					9960		
US	6245771			В1	2	2001()612	US	3]	1998-	2995	1		19	9980:	317	<

US 6274586 B1 20010814 US 2000-571402 20000515 <-PRIORITY APPLN. INFO.:

DE 1995-19535516 A 19950925

WO 1996-EP4013 W 19960912
US 1998-29951 A3 19980317

OTHER SOURCE(S): MARPAT 126:273648

AB The invention relates to means of combating parasitic fungi containing as the active agents at least one compound which prevents respiration in the cytochrome complex III and fenazaquin. The invention may be used in particular in combating Botrytis.

IT 189005-47-8D, mixts. with fenazaquin

RL: AGR (Agricultural use); BIOL (Biological study); USES (Uses) (synergistic agrochem. fungicides)

RN 189005-47-8 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -(methoxyimino)-N-methyl- (9CI) (CA INDEX NAME)

L89 ANSWER 51 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1997:275760 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 127:5034

TITLE: An improved general method for the preparation of

4-aryl substituted bispyrazolo[3,4-b;4',3'-e]pyridines

AUTHOR(S): Puchala, Agnieszka; Rasala, Danuta; Kolehmainen,

Erkki; Prokesova, Monika

CORPORATE SOURCE: Institute of Chemistry, Pedagogical University,

Kielce, PL-25-020, Pol.

SOURCE: Organic Preparations and Procedures International (

1997), 29(2), 226-230

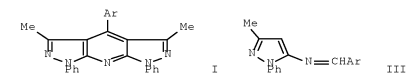
CODEN: OPPIAK; ISSN: 0030-4948

PUBLISHER: Organic Preparations and Procedures, Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 127:5034

GΙ



AB The title compds. I (Ar = Ph, substituted Ph, 2-furoyl, 4-pyridyl) were prepared by reacting 5-amino-3-methyl-1-phenylpyrazole (II) with ArCHO or via reaction of Schiff bases III with II.

IT 186140-69-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of aryl bispyrazolopyridines)

RN 186140-69-2 ZCAPLUS

CN Benzoic acid, 4-[[(3-methyl-1-phenyl-1H-pyrazol-5-yl)imino]methyl]-, methyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT: 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L89 ANSWER 52 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1997:14891 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 126:46975

TITLE: Preparation of (hetero)aryloxycrotonates and related

compounds as insecticides and fungicides.

INVENTOR(S): Grote, Thomas; Kirstgen, Reinhard; Mueller, Bernd;

Sauter, Hubert; Harreus, Albrecht; Koenig, Hartmann; Ammermann, Eberhard; Lorenz, Gisela; Strathmann,

Siegfried; Roehl, Franz

PATENT ASSIGNEE(S): BASF A.-G., Germany

SOURCE: PCT Int. Appl., 219 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	TENT NO.	KINI	D DATE	APPLICATION NO.	DATE
CA AU	W: AU, BG, UA, US, RW: AT, BE, 2217773 9656483	BR, CA, AM, AZ, CH, DE, A1 A	CN, CZ, HU, BY, KG, KZ, DK, ES, FI, 19961114 19961129	WO 1996-EP1754 JP, KR, MX, NO, NZ, MD, RU, TJ, TM FR, GB, GR, IE, IT, CA 1996-2217773 AU 1996-56483	PL, SG, SK, TR, LU, MC, NL, PT, SE 19960426 < 19960426 <
	824518 824518		20010627	EP 1996-913530	
HU BR JP AT ZA US	1187814	A A2 A T T A A			19960426 < 19960426 < 19960426 < 19960426 < 19960426 < 19960508 < 19971030 <

OTHER SOURCE(S):

MARPAT 126:46975

GΙ

$$\begin{array}{c|c}
 & \text{Y} & \text{Z} \\
 & \text{X} & \text{V} & \text{Cour}^1 \\
 & \text{C} & \text{CHR}^2 & \text{I}
\end{array}$$

Title compds. (I; U = O, S, NH; V = O, S, NH, alkylimino; X, Y, Z = N, CR3; R1, R2 = alkyl; R3 = H, cyano, NO2, halo, alkyl, haloalkyl, alkoxy, haloalkoxy, alkylthio haloalkylthio; R4 = organic group bound directly or via O, S, imino, carboxyl, or CONH), were prepared as insecticides and agrochem. fungicides (no data). Thus, to a solution of KOH in DMF was added 3-iodophenol and then Me 3-bromocrotonate; the mixture was stirred 1 h at room temperature to give 61% Me α -(3-iodophenoxy)crotonate. The latter was refluxed with 4-chlorophenylboronic acid and Pd(Ph3)4 in H2O/dimethoxyethane to give 90% Me 2-(4-chlorobiphenyl-4-yloxy)but-2- enoate.

IT 184883-56-5P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of (hetero)aryloxycrotonates and related compds. as insecticides and fungicides)

RN 184883-56-5 ZCAPLUS

CN 2-Butenoic acid, 2-[3-[2-[4-chloro-3-(4-chlorophenyl)-5-isoxazolyl]ethenyl]phenoxy]-, methyl ester (9CI) (CA INDEX NAME)

L89 ANSWER 53 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1997:4342 ZCAPLUS Full-text

DOCUMENT NUMBER: 126:74851

TITLE: Preparation of azolyloxybenzylalkoxyacrylates as

agrochemical fungicides.

INVENTOR(S): Mueller, Bernd; Kirstgen, Reinhard; Koenig, Hartmann;

Rack, Michael; Oberdorf, Klaus; Roehl, Franz; Sauter,

Hubert; Lorenz, Gisela; Ammermann, Eberhard

PATENT ASSIGNEE(S): BASF A.-G., Germany

SOURCE: Ger. Offen., 34 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Faient LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	TENT NO.															
DE	19519041			A1		1996	1128	DE	1995-	-1951	9041		1			
IL	118168			Α		2001	0724	IL	1996-	-1181	68		1	9960	507	<
CA	118168 2218897			A1		1996	1128	CA	1996-	-2218	897		1	9960		
WO	9637477			A1		1996	1128	WO	1996-	-EP20	42		1			
	W: AU,															
								MD, RI			·	·	ĺ	·	·	
	RW: AT,	BE,	CH,	DE,	DK,	ES,	FΙ,	FR, G	B, GR	, IE,	IT,	LU,	MC,	NL,	PT,	SE
AU	9658956			Α		1996	1211	AU	1996-	-5895	6		1	9960!	513	<
AU	712768			В2		1999	1118									
EP	830342			A1		1998	0325	EP	1996-	-9160	55		1	9960!	513	<
EP	830342 830342			В1		2002	1009									
	R: AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB, G	R, IT,	LI,	NL,	SE,	PT,	IE		
CN	1185148			Α		1998	0617	CN	1996-	-1941	36		1	9960	513	<
CN	1069638			В		2001	0815									
								BR								
JP	11511744															
								NZ						9960	513	<
EP	1110453					2001	0627	EP	2001-	-1076	39		1	9960	513	<
EP	1110453			В1		2003	0502									
	R: AT,															
AT	225773 830342			Τ		2002	1015	AT	1996-	-9160	55		1	9960	513	
PT	830342			Τ		2003	0228	PT	1996-	-9160	55		1	9960	513	
AT	238660			Τ		2003	0515	AT	2001-	-1076	39		1	.9960!	513	
ES	2187653								1996-	-9160	55		1	9960	513	
									1996-							
								US	1997-	-9527	55		1			
	6380231			В1		2002	0430	US	1999-	-2872	74		1	9990		<
PRIORITY	Y APPLN.	INFO.	:						1995-							
								EP	1996-	-9160	55		A3 1	9960!	513	
								WO	1996-	-EP20	42	,	W 1	.9960!	513	
									1997-	-9527	55		A3 1	9971	120	
OTHER SO	OURCE(S):			MARE	PAT	126:	74851	1								

$$CO_2R1$$
 $CHOR^2$
 OMe
 OMe

AB Title compds. [I; n = 0-4; R = NO2, cyano, halo, (substituted) alkyl, alkenyl, alkynyl, alkoxy, alkenyloxy; adjacent R groups may form a bridge; R1, R2 = alkyl; R3 = substituted pyrazolyl, triazolyl], were prepared Thus, Me α -(2-bromomethylphenyl)- β -methoxyacrylate and 1-(o-chlorophenyl)-3-hydroxy-

1,2,4-triazole were stirred with Na2CO3 in DMF to give 12% title compound (II). II at 63 ppm on wheat seedlings reduced incidence of Puccinia recondita to \leq 15% vs. 65% for untreated controls.

IT 184684-07-9P 184684-08-0P 184684-09-1P 184684-10-4P 184684-11-5P 184684-12-6P 184684-13-7P 184684-15-9P 184684-22-8P 184684-23-9P 184684-24-0P 184684-25-1P 184684-26-2P 184684-27-3P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of azolyloxybenzylalkoxyacrylates as agrochem. fungicides)

RN 184684-07-9 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(4-\text{chlorophenyl})-1H-\text{pyrazol}-3-\text{yl}] \text{oxy}] \text{methyl}] - \alpha-(\text{methoxymethylene})-, \text{methyl ester (9CI)}$ (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 184684-08-0 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[[[1-(4-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 184684-09-1 ZCAPLUS

CN Benzeneacetic acid, 2-chloro-6-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -(methoxymethylene)-, methyl ester (9CI) (CA INDEX NAME)

RN 184684-10-4 ZCAPLUS

CN Benzeneacetic acid, 2-chloro-6-[[[1-(2,4-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -(methoxymethylene)-, methyl ester (9CI) (CA INDEX

NAME)

RN 184684-11-5 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[[[1-[5-(trifluoromethyl)-2-pyridinyl]-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 184684-12-6 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[[[1-[4-(trifluoromethoxy)phenyl]-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 184684-13-7 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[[(1-pyrazinyl-1H-pyrazol-3-yl)oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} N & N & O-CH_2 \\ \hline \\ N & MeO-C-C \\ \hline \\ M-OMe \\ \end{array}$$

RN 184684-15-9 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(6-chloro-2-pyridinyl)-1H-pyrazol-3-yl]oxy]methyl]- α -(methoxymethylene)-, methyl ester (9CI) (CA INDEX NAME)

RN 184684-22-8 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(4-chlorophenyl)-1H-pyrazol-4-yl]oxy]methyl]- α -(methoxymethylene)-, methyl ester (9CI) (CA INDEX NAME)

RN 184684-23-9 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(2-chloro-4-fluorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -(methoxymethylene)-, methyl ester (9CI) (CA INDEX NAME)

RN 184684-24-0 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(2,4-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -(methoxymethylene)-, methyl ester (9CI) (CA INDEX

NAME)

RN 184684-25-1 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(5-chloro-2-pyridinyl)-1H-pyrazol-3-yl]oxy]methyl]- α -(methoxymethylene)-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} C1 & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

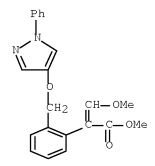
RN 184684-26-2 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(2,4-difluorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -(methoxymethylene)-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{CH-OMe} & \text{O} \\ \hline \\ \text{CH}_2 - \text{O} \\ \hline \end{array}$$

RN 184684-27-3 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[[(1-phenyl-1H-pyrazol-4-yl)oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)



L89 ANSWER 54 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1996:718922 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 126:117690

TITLE: 1-Phenyl-3-methyl-5-N-benzylideneaminopyrazoles.

Substituent effects and protonation sites studied by

NMR and ab initio (6-31G*) MO calculations

AUTHOR(S): Kolehmainen, Erkki; Puchala, Agnieszka; Suontamo,

Reijo; Rasala, Danuta; Lysek, Robert

CORPORATE SOURCE: Dep. Chem., Univ. Jyvaskyla, Jyvaskyla, FIN-40351,

Finland

SOURCE: Journal of the Chemical Society, Perkin Transactions

2: Physical Organic Chemistry (1996), (11),

2383-2387

CODEN: JCPKBH; ISSN: 0300-9580 Royal Society of Chemistry

DOCUMENT TYPE: Journal LANGUAGE: English

1-Phenyl-3-methyl-5-N-benzylideneaminopyrazole and its derivs. 11 prepared by AΒ condensation of 1-phenyl-3-methyl-5-aminopyrazole and aromatic aldehydes have been studied by multinuclear (1H, 13C, 14/15N and 170) magnetic resonance spectroscopy. The 13C NMR chemical shifts and the direct spin-spin coupling consts. 1J(C,H) of the azomethine carbon of these Schiff bases (SB) correlate significantly with the Hammett substituent consts., σp , of the parasubstituents in the aryl ring bound to the azomethine carbon. The assignments of the 15N NMR chems. shifts of SBs in CDC13 were based on 2J(N,H)s observed for the azomethine nitrogen as well as 1H, 15N HMBC expts. Based on the present 1H, 13C and 15N NMR data these SBs can be transformed to single and double protonated forms in trifluoroacetic acid (TFA). The protonation sites (the first on e at the unsubstituted nitrogen of the pyrazole ring and the second one at the azomethine nitrogen) deduced from the NMR data are supported by ab initio MO calcns. at $HF/6-31G^*$ level with a full geometry optimization performed for a model compound, 1,3-dimethyl-5-N-benzylideneaminopyrazole.

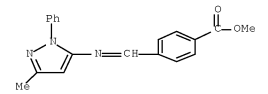
IT 186140-69-2P

PUBLISHER:

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (multinuclear magnetic resonance of Schiff bases)

RN 186140-69-2 ZCAPLUS

CN Benzoic acid, 4-[[(3-methyl-1-phenyl-1H-pyrazol-5-yl)imino]methyl]-, methyl ester (9CI) (CA INDEX NAME)



L89 ANSWER 55 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1996:457757 ZCAPLUS Full-text

DOCUMENT NUMBER: 125:114606

TITLE: Preparation of (pyrazolylmethyl)thiazolidines useful

as hypoglycemic agents and aldose-reductase inhibitors

INVENTOR(S): Ohara, Yoshio; Suzuki, Mikio; Miyachi, Nobuhide; Kato,

Katsuhiro; Ohdoi, Keisuke; Kobayashi, Tetsuya;

Shikada, Ken-ichi; Naito, Takeshi; Yotsumoto, Takashi

PATENT ASSIGNEE(S): Nissan Chemical Industries, Ltd., Japan

SOURCE: PCT Int. Appl., 209 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
WO 9611196	A1 19960418	WO 1995-JP2041	19951005 <
W: AU, CA, CN,	CZ, FI, HU, KR, L	T, MX, NO, NZ, RO, RU	, SI, SK, UA, US
RW: AT, BE, CH,	DE, DK, ES, FR, GI	B, GR, IE, IT, LU, MC	, NL, PT, SE
JP 08157473	A 19960618	JP 1995-246171	19950925 <
AU 9536190	A 19960502	AU 1995-36190	19951005 <
ZA 9508395	A 19960514	ZA 1995-8395	19951005 <
PRIORITY APPLN. INFO.:		JP 1994-242865	A 19941006
		JP 1995-246171	A 19950925
		WO 1995-JP2041	W 19951005

OTHER SOURCE(S): MARPAT 125:114606

GΙ

$$\begin{array}{c|c}
 & R3 \\
 & X2 \\
 & X2
\end{array}$$

$$\begin{array}{c|c}
 & R4 & O \\
 & NR5 \\
 & X2
\end{array}$$

AB The title compds [I; X1 = S, O; X2 = S, O, NH; Y = C(R6)R7; R6, R7 = H, alkyl, cycloalkyl; R1 = alkyl, alkoxy, etc.; R2, R3 = H, alkyl, etc.; R4 = H, alkyl, etc.; R5 = H, CO2Me], useful as antidiabetic agents and aldose-reductase inhibitors for the treatment of diabetes mellitus and its complications, are prepared and I-containing formulations presented. Thus, 5-[[5-(2-hydroxy-2-phenylethoxy)-1-methyl-3-pyrazolyl]methylidene]thiazoli din-2,4-dione,

prepared in a multiple-step procedure from Et -5-hydroxy-1-methyl-3-pyrazolecarboxylate, demonstrated a 42.3% anti-glycation effect as determined by the Lowry method at 0.24 mM.

IT 179099-21-9p 179099-22-0p 179099-26-4p 179099-29-7p

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of (pyrazolylmethyl)thiazolidines useful as hypoglycemic agents

and aldose-reductase inhibitors)

RN 179099-21-9 ZCAPLUS

CN 1H-Pyrazole-3-carboxylic acid, 1-methyl-5-[2-(5-methyl-2-phenyl-4-oxazolyl)-2-oxoethoxy]-, ethyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{Ph} & \overset{\text{N}}{\longrightarrow} & \overset{\text{N}}{\longleftarrow} & \overset{\text{Me}}{\longrightarrow} & \overset{\text{N}}{\longrightarrow} & \overset{N}{\longrightarrow} & \overset{\text{N}}{\longrightarrow} & \overset{\text{N}}{\longrightarrow} & \overset{\text{N}}{\longrightarrow} & \overset{\text{N}}{\longrightarrow} & \overset{\text{N$$

RN 179099-22-0 ZCAPLUS

CN 1H-Pyrazole-3-carboxylic acid, 1-(1,1-dimethylethyl)-5-[2-(5-methyl-2-phenyl-4-oxazolyl)-2-oxoethoxy]-, ethyl ester (9CI) (CA INDEX NAME)

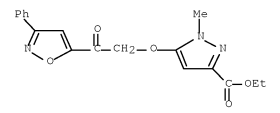
$$\begin{array}{c} \text{Ph} & \overset{\circ}{\underset{\text{Me}}{\bigvee}} \text{CH}_2 - \overset{\text{t-Bu}}{\underset{\text{N}}{\bigvee}} \text{N} \\ & \overset{\circ}{\underset{\text{N}}{\bigvee}} \text{C-OEt} \end{array}$$

RN 179099-26-4 ZCAPLUS

CN 1H-Pyrazole-3-carboxylic acid, 1-methyl-5-[2-(5-methyl-1-phenyl-1H-pyrazol-4-yl)-2-oxoethoxy]-, ethyl ester (9CI) (CA INDEX NAME)

RN 179099-29-7 ZCAPLUS

CN 1H-Pyrazole-3-carboxylic acid, 1-methyl-5-[2-oxo-2-(3-phenyl-5-isoxazolyl)ethoxy]-, ethyl ester (9CI) (CA INDEX NAME)



L89 ANSWER 56 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1996:410459 ZCAPLUS Full-text

DOCUMENT NUMBER: 125:86315

TITLE: Preparation of alkyl phenylacetate pesticides and

agrochemical fungicides

INVENTOR(S): Oberdorf, Klaus; Sauter, Hubert; Koenig, Hartmann;

Harreus, Albrecht; Mueller, Bernd; Kirstgen, Reinhard; Grammenos, Wassilios; Bayer, Herbert; Roehl, Franz; et

al.

PATENT ASSIGNEE(S): BASF A.-G., Germany SOURCE: PCT Int. Appl., 561 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Fatent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA.	TENT 1	NO.			KINI)	DATE		P	APPL:	ICAT	ION 1	NO.		D	ATE		
WO	9607	 633			A1	_	1996	0314	_ ⊽	VO 1	 995-:	EP34	05		1	9950	830	<
	W:	ΑU,	BG,	BR,	BY,	CA,	CN,	CZ,	FI,	HU,	JP,	KR,	KΖ,	MX,	NO,	NZ,	PL,	
		RU,	SG,	SK,	UA,	US												
	RW:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE	
CA	2199	422			A1		1996	0314		CA 1	995-	2199	422		1	9950	830	<
AU	9533	878			Α		1996	0327	I	AU 19	995-	3387	8		1	9950	830	<
EP	7812	66			A1		1997	0702	E	EP 19	995-	9305.	31		1	9950	830	<
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	IT,	LI,	NL,	PT,	SE		
CN	1161	687			Α		1997	1008		CN 19	995-	1958.	27		1	9950	830	<
BR	9509	004			Α		1998	0602	E	BR 1	995-	9004			1	9950	830	<
JP	1050	5596			Τ		1998	0602	J	JP 1	995-	5091	72		1	9950	830	<
ZA	9507	545			Α		1997	0310	2	ZA 19	995-	7545			1	9950	908	<
PRIORIT	Y APP	LN.	INFO	.:					Γ	DE 1	994-	4432	336		A 1	9940	910	
									V	VO 1	995-	EP34	05		W 1	9950	830	
OTHED C	OLIDOH.	(C).			MADI	ייי ער	105.	0621	=									

OTHER SOURCE(S): MARPAT 125:86315

GI

- AB The title compds. [I; R = halogen, hydroxy, mercapto, amino, carboxyl, carbonylamino, etc.; R1 = CHO, alkylcarbonyl, alkyl; R2 = alkyl; U = O, S, NH, NHO; V = O, S, NH; X = CN, NO2, halogen, (halo)alkyl, (halo)alkoxy, alkylthio, etc.; n = 0-3], useful as agrochem. fungicides and pesticides, are prepared Thus, Me α -[2-(2-methylphenoxymethylene)phenyl]- α ketoacetate was reacted with NaBH4 and HCl, , and the intermediate treated with NaH and MeI, producing pesticidal phenylacetate ester II.

RL: AGR (Agricultural use); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of alkyl phenylacetate pesticides and agrochem. fungicides)

RN 178428-10-9 ZCAPLUS

CN Benzeneacetic acid, α -methoxy-2-[[(1-phenyl-1H-pyrazol-3-yl)oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

- RN 178428-11-0 ZCAPLUS
- CN Benzeneacetic acid, α -methoxy-2-[[[1-(4-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

- RN 178428-12-1 ZCAPLUS
- CN Benzeneacetic acid, $2-[[[1-(4-\text{chlorophenyl})-1H-\text{pyrazol}-3-\text{yl}] \text{oxy}] \text{methyl}]-\alpha-\text{methoxy-, methyl ester (9CI)}$ (CA INDEX NAME)

RN 178428-13-2 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(2,4-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -methoxy-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\$$

RN 178428-19-8 ZCAPLUS

CN Benzeneacetic acid, α -methoxy-2-[[[1-[5-(trifluoromethyl)-2-pyridinyl]-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 178428-20-1 ZCAPLUS

CN Benzeneacetic acid, 2-[[[4-chloro-1-[5-(trifluoromethyl)-2-pyridinyl]-1H-pyrazol-3-yl]oxy]methyl]- α -methoxy-, methyl ester (9CI) (CA INDEX NAME)

CN Benzeneacetamide, 2-[[[1-(2,4-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -methoxy-N-methyl- (9CI) (CA INDEX NAME)

RN 178428-66-5 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -methoxy-N-methyl- (9CI) (CA INDEX NAME)

RN 178428-71-2 ZCAPLUS

CN Benzeneacetamide, α -methoxy-N-methyl-2-[[(1-phenyl-1H-pyrazol-3-yl)oxy]methyl]- (9CI) (CA INDEX NAME)

RN 178428-72-3 ZCAPLUS

CN Benzeneacetamide, α -methoxy-N-methyl-2-[[[1-(4-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]- (9CI) (CA INDEX NAME)

RN 178428-87-0 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]-

 α -ethoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 178428-94-9 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(4-fluorophenyl)-1H-pyrazol-3-yl]oxy]methyl]-\alpha-methoxy-, methyl ester (9CI) (CA INDEX NAME)$

RN 178428-95-0 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(4-fluorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -methoxy-N-methyl- (9CI) (CA INDEX NAME)

RN 178428-96-1 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -ethoxy-N-methyl- (9CI) (CA INDEX NAME)

L89 ANSWER 57 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1996:399852 ZCAPLUS Full-text

DOCUMENT NUMBER: 125:184056

TITLE: Synthesis and complexation of macrocycles containing

two pyrazolone sub-units

AUTHOR(S): Marzin, C.; Naji, M.; Coquelet, C.; Tarrago, G. CORPORATE SOURCE: Equipe Chimie Supramoleculaire, LMPM, UMR 5635,

Equipe Chimie Supramoleculaire, LMPM, UMR 5635, Universite Montpellier II, Montpellier, 34095, Fr.

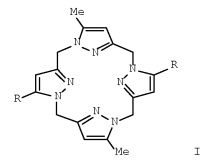
SOURCE: Inorganica Chimica Acta (1996), 246(1-2),

217-227

CODEN: ICHAA3; ISSN: 0020-1693

PUBLISHER: Elsevier
DOCUMENT TYPE: Journal
LANGUAGE: English

GΙ



The synthesis and characterization of several Ru(II) complexes with acyclic and macrocyclic ligands containing tautomerizable OH and fixed OCH3 5-pyrazolone heterocycles are described. From dipyrazolylmethane bidentate ligands L, RuL(bpy)2(PF6)2 and Ru(L-H+)(bpy)2PF6 complexes were obtained. From the macrocycle with two CH3 and two OCH3 pyrazole sub-units (I; R = OMe), Ru(I)XY(PF6)2 (X, Y = DMSO, MeCN, Py, pyrazole, 3,5-dimethylpyrazole) were prepared They show a behavior close to that of the analogous tetrapyrazole complexes but with slightly different complexing ability. In the case of I (R = OH), coordination with Ru(DMSO)4C12 leads to unstable complexes.

IT 180518-76-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(for preparation of pyrazole derivs. or pyrazole-based macrocycles and their

ruthenium complexes)

RN 180518-76-7 ZCAPLUS

CN 1H-Pyrazole-3-carboxylic acid, 5-methoxy-1-[[5-methyl-1-(tetrahydro-2H-pyran-2-yl)-1H-pyrazol-3-yl]methyl]-, ethyl ester (9CI) (CA INDEX NAME)

L89 ANSWER 58 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1996:231375 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 124:261031

TITLE: Preparation of [2-(pyrazolylvinyl)phenyl]methoximino-N-

methylacetamides as pesticides

INVENTOR(S): Kirstgen, Reinhard; Koenig, Hartmann; Sauter, Hubert;

Harries, Volker; Lorenz, Giesela; Ammermann, Eberhard

PATENT ASSIGNEE(S): BASF A.-G., Germany SOURCE: Eur. Pat. Appl., 26 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Fatent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.					KIND		DATE		A	APPLICATION NO.						DATE	
	EP	691332	?			A1	_	1996	0110	E:	 Р 1	.995–	1099	81		-	19950627	<
	ΕP	691332	2			В1		1999	0908									
		R: <i>I</i>	ΔT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	ΙT,	LI,	NL,	PΊ	S, SE	
	ΑT	184276	-)			T		1999	0915	А	T 1	995-	1099	81			19950627	<
	ES	213741	. 1			Т3		1999	1216	E	S 1	995-	1099	81			19950627	<
	CA	215299	6			A1		1996	0107	C.	A 1	995-	2152	2996			19950629	<
	JΡ	080534	120			Α		1996	0227	J.	P 1	995-	1638	348			19950629	<
	AU	952482	8.2			Α		1996	0118	A	U 1	995-	2482	28			19950704	<
	ΑU	684640)			В2		1997	1218									
	US	550625	54			Α		1996	0409	U	S 1	995-	4987	759			19950706	<
	CN	112233	30			Α		1996	0515	C	N 1	995-	1083	316			19950706	<
PRIOR	YTI?	APPLN	1.]	INFO	. :					D	E 1	994-	4423	615		А	19940706	
OTHER	OTHER SOURCE(S):				MARI	PAT	124:	26103	31									
GI																		

AB Title compds. [I; R = CR4:CHZC(:NOMe)CONHMe; R2 = H, alkyl, heterocyclyl, (hetero)aryl, etc.; R3 = cyano, (halo)alkyl, alkoxy, etc.; R4 = H, cyano, halo, (halo)alkyl; Z = (un)substituted 1,2-phenylene; m = 0-2] were prepared Thus, 2-[(MeO)2P(O)CH2]C6H4C(:NOMe)CONHMe (preparation given) was condensed with 1-(2,4-dichlorophenyl)-4-formyl-5-methylpyrazole to give title compound (E,E)-II which gave ≥85% control of Paricularia oryzae on rice seedlings at 250ppm.

IT 175424-53-0P 175424-54-1P 175424-55-2P 175424-56-3P 175424-57-4P 175424-58-5P 175424-60-9P 175424-61-0P 175424-62-1P 175424-63-2P 175424-64-3P 175424-65-4P 175424-66-5P 175424-67-6P 175424-68-7P

175424-69-8P 175424-70-1P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of [2-(pyrazolylvinyl)phenyl]methoximino-N-methylacetamides as pesticides)

RN 175424-53-0 ZCAPLUS

CN Benzeneacetamide, 2-[2-[1-(2,4-dichlorophenyl)-5-(trifluoromethyl)-1H-pyrazol-4-yl]ethenyl]- α -(methoxyimino)-N-methyl-, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 175424-54-1 ZCAPLUS

CN Benzeneacetamide, 2-[2-[1-(3-chlorophenyl)-5-(trifluoromethyl)-1H-pyrazol-4-yl]ethenyl]- α -(methoxyimino)-N-methyl-, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 175424-55-2 ZCAPLUS

CN Benzeneacetamide, α -(methoxyimino)-N-methyl-2-[2-[1-(4-methylphenyl)-5-(trifluoromethyl)-1H-pyrazol-4-yl]ethenyl]-, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 175424-56-3 ZCAPLUS

CN Benzeneacetamide, 2-[2-[1-(4-chlorophenyl)-5-(trifluoromethyl)-1H-pyrazol-4-yl]ethenyl]- α -(methoxyimino)-N-methyl-, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 175424-57-4 ZCAPLUS

CN Benzeneacetamide, 2-[2-[1-(2,4-dichlorophenyl)-1H-pyrazol-4-yl]ethenyl]- $\alpha-(methoxyimino)-N-methyl-$, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 175424-58-5 ZCAPLUS

CN Benzeneacetamide, $2-[2-[1-(4-\text{chlorophenyl})-1H-\text{pyrazol}-4-\text{yl}]\text{ethenyl}]-\alpha-(\text{methoxyimino})-N-\text{methyl}-, (E,E)- (9CI) (CA INDEX NAME)$

Double bond geometry as shown.

RN 175424-60-9 ZCAPLUS

CN Benzeneacetamide, 2-[2-[1-(2,4-dichlorophenyl)-5-methyl-1H-pyrazol-4-yl]ethenyl]- α -(methoxyimino)-N-methyl-, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 175424-61-0 ZCAPLUS

CN Benzeneacetamide, 2-[2-[5-chloro-1-(2,4-dichlorophenyl)-1H-pyrazol-4-yl]ethenyl]- α -(methoxyimino)-N-methyl-, (E,E)- (9CI) (CA INDEX NAME)

RN 175424-62-1 ZCAPLUS

CN Benzeneacetamide, 2-[2-[5-chloro-1-(4-chlorophenyl)-1H-pyrazol-4-yl]ethenyl]- α -(methoxyimino)-N-methyl-, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 175424-63-2 ZCAPLUS

CN Benzeneacetamide, 2-[2-[5-chloro-1-(3-chlorophenyl)-1H-pyrazol-4-yl]ethenyl]- α -(methoxyimino)-N-methyl-, (E,E)- (9CI) (CA INDEX NAME)

RN 175424-64-3 ZCAPLUS

CN Benzeneacetamide, α -(methoxyimino)-N-methyl-2-[2-(1-phenyl-1H-pyrazol-4-yl)ethenyl]-, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 175424-65-4 ZCAPLUS

CN Benzeneacetamide, 2-[2-[1-(2-chlorophenyl)-5-(trifluoromethyl)-1H-pyrazol-4-yl]ethenyl]- α -(methoxyimino)-N-methyl-, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 175424-66-5 ZCAPLUS

CN Benzeneacetamide, 2-[2-[1-(4-fluorophenyl)-5-(trifluoromethyl)-1H-pyrazol-4-yl]ethenyl]- α -(methoxyimino)-N-methyl-, (E,E)- (9CI) (CA INDEX NAME)

RN 175424-67-6 ZCAPLUS

CN Benzeneacetamide, 2-[2-[1-(3,5-dichlorophenyl)-5-(trifluoromethyl)-1H-pyrazol-4-yl]ethenyl]- α -(methoxyimino)-N-methyl-, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 175424-68-7 ZCAPLUS

CN Benzeneacetamide, α -(methoxyimino)-2-[2-[1-(4-methoxyphenyl)-5-(trifluoromethyl)-1H-pyrazol-4-yl]ethenyl]-N-methyl-, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 175424-69-8 ZCAPLUS

CN Benzeneacetamide, α -(methoxyimino)-N-methyl-2-[2-[1-(3-methylphenyl)-5-(trifluoromethyl)-1H-pyrazol-4-yl]ethenyl]-, (E,E)- (9CI) (CA INDEX NAME)

RN 175424-70-1 ZCAPLUS

CN Benzeneacetamide, $2-[2-[1-(3-\text{chlorophenyl})-1H-\text{pyrazol}-4-\text{yl}] = thenyl] - \alpha-(methoxyimino)-N-methyl-, (E,E)- (9CI) (CA INDEX NAME)$

Double bond geometry as shown.

L89 ANSWER 59 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1996:205034 ZCAPLUS Full-text

DOCUMENT NUMBER: 124:261025

TITLE: Preparation of N-methoxy-N-

[(pyrazolyloxymethyl)phenyl]carbamates and analogs as

agrochemical fungicides and pesticides

INVENTOR(S): Mueller, Bernd; Koenig, Hartmann; Kirstgen, Reinhard;

Oberdorf, Klaus; Roehl, Franz; Goetz, Norbert; Sauter,

Hubert; Lorenz, Gisela; Ammermann, Eberhard

PATENT ASSIGNEE(S): BASF A.-G., Germany SOURCE: Ger. Offen., 47 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Fatent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 4423612	A1	19960111	DE 1994-4423612	19940706 <
CA 2194503	A1	19960118	CA 1995-2194503	19950621 <
CA 2194503	С	20070424		
WO 9601256	A1	19960118	WO 1995-EP2396	19950621 <

W: AU, BG, BR, BY, CA, CN, CZ, FI, HU, JP, KR, KZ, MX, NO, NZ, PL, RU, SG, SK, UA, US RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE AU 9529222 19960125 AU 1995-29222 19950621 <--Α AU 685299 19980115 В2 CN 1154692 19970716 CN 1995-194436 19950621 <--Α CN 1068313 В 20010711 BR 9508242 19970930 BR 1995-8242 19950621 <--Α EP 1995-924888 EP 804421 Α1 19971105 19950621 <--EP 804421 В1 19980916 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, NL, SE, PT, IE Τ 19980512 JP 1996-503648 19950621 <--JP 10504810 JP 3838659 В2 20061025 HU 77510 A2 19980528 HU 1997-29 19950621 <--HU 218298 20000728 R AT 171165 Τ 19981015 AT 1995-924888 19950621 <--ES 2123264 Т3 19990101 ES 1995-924888 19950621 <--19950621 <--RU 2151142 C1 20000620 RU 1997-102108 PL 180298 PL 1995-318100 19950621 <--В1 20010131 SK 282426 В6 20020107 SK 1997-17 19950621 <--PL 186501 В1 20040130 PL 1995-340891 19950621 CZ 294484 20050112 CZ 1997-37 В6 19950621 IL 114390 20010128 IL 1995-114390 19950629 <--Α ZA 9510727 19970618 ZA 1995-10727 19951218 <--Α NO 9700042 NO 1997-42 19970305 19970106 <--Α NO 307336 20000320 В1 US 5869517 19990209 US 1997-765185 19970106 <--Α FI 9700067 19970305 FI 1997-67 19970107 <--Α FI 117199 В1 20060731 BG 63081 В1 20010330 BG 1997-101198 19970204 <--US 6054592 20000425 US 1998-131640 19980810 <--Α CN 1308065 20010815 CN 2000-129025 20000927 <--Α PRIORITY APPLN. INFO.: DE 1994-4423612 19940706 WO 1995-EP2396 19950621 OTHER SOURCE(S): CASREACT 124:261025; MARPAT 124:261025

 $Q = R3N \longrightarrow 0 \qquad R3N \longrightarrow 0 \qquad X1Me$ $R3N \longrightarrow 0 \longrightarrow X1Me$

GΙ

AB RCH2ZN(OR4)COZ1R5 [R = pyrazolyloxy group Q; R2 = halo, alkyl, alkoxy, etc.; R3 = alk(en)yl, heterocyclyl, (hetero)aryl, etc.; R4 = H, alkyl, alkanoyl, alkoxycarbonyl, etc.; R5 = H, (cyclo)alk(en)yl, alkynyl; Z = (un)substituted 1,2-phenylene; Z1 = bond, O, (alkyl)imino, etc.; m = 0-2] were prepared Thus, 2-MeC6H4NHOH was amidated by C1CO2Ph and the product converted in 2 steps to give 2-(BrCH2)C6H4N(OMe)CO2Ph which was condensed with N-(2-pyrazinyl)-3-hydroxypyrazole to give, after NHMe amidation, title compound II (R3 = 2-pyrazinyl, Z1 = NH). II (R3 = 4-C1C6H4, Z1 = 0) gave ≥95% control of Puccinia recondita on wheat seedlings at 63ppm.

IT 175013-18-0P 175013-19-1P 175013-20-4P

175013-21-5P 175013-22-6P 175013-23-7P 175013-24-8P 175013-25-9P 175013-26-0P 175013-27-1P 175013-28-2P 175013-29-3P 175013-30-6P 175013-31-7P 175013-33-9P 175013-34-0P 175013-35-1P 175013-36-2P 175013-37-3P 175013-38-4P 175013-39-5P 175013-40-8P 175013-42-0P 175013-43-1P RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of N-methoxy-N-[(pyrazolyloxymethyl)phenyl]carbamates and analogs as agrochem. fungicides and pesticides) 175013-18-0 ZCAPLUS RN Carbamic acid, N-[2-[[[1-(4-chlorophenyl)-1H-pyrazol-3-CMyl]oxy]methyl]phenyl]-N-methoxy-, methyl ester (CA INDEX NAME)

RN 175013-19-1 ZCAPLUS
CN Carbamic acid, [2-[[[1-(2,4-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-20-4 ZCAPLUS
CN Carbamic acid, methoxy[2-[[[1-(2-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-21-5 ZCAPLUS
CN Carbamic acid, methoxy[2-[[[1-(3-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-22-6 ZCAPLUS

CN Carbamic acid, methoxy[2-[[[1-(4-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-23-7 ZCAPLUS

CN Carbamic acid, [2-[[[1-(2-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-24-8 ZCAPLUS

CN Carbamic acid, [2-[[[1-(3-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{Cl} & \text{N} & \text{N} & \text{O-CH}_2 \\ \text{MeO-C-N} & \text{OMe} \end{array}$$

RN 175013-25-9 ZCAPLUS

CN Carbamic acid, [2-[[[1-(2,6-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-26-0 ZCAPLUS

CN Carbamic acid, [2-[[[1-(3,5-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-27-1 ZCAPLUS

CN Carbamic acid, [2-[[[1-(2,5-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & &$$

RN 175013-28-2 ZCAPLUS

CN Carbamic acid, [2-[[[1-(3,4-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-29-3 ZCAPLUS

CN Carbamic acid, [2-[[[1-(4-chloro-2-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-30-6 ZCAPLUS

CN Carbamic acid, methoxy[2-[[[1-[3-(trifluoromethyl)phenyl]-1H-pyrazol-3-yl]oxy]methyl]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-31-7 ZCAPLUS

CN Carbamic acid, methoxy[2-[[[1-(4-methoxyphenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-33-9 ZCAPLUS

CN Carbamic acid, [2-[[[1-(4-fluorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 175013-34-0 ZCAPLUS

CN Carbamic acid, [2-[[[4-chloro-1-(4-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-35-1 ZCAPLUS

CN Carbamic acid, methoxy[2-[[[1-(3-methoxyphenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-36-2 ZCAPLUS

CN Carbamic acid, [2-[[[4-chloro-1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

RN 175013-37-3 ZCAPLUS

CN Carbamic acid, [2-[[[1-(2,2-difluoro-1,3-benzodioxol-5-yl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]methoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-38-4 ZCAPLUS

CN Carbamic acid, methoxy[2-[[[1-(2-pyridinyl)-1H-pyrazol-3-yl]oxy]methyl]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-39-5 ZCAPLUS

CN Carbamic acid, methoxy[2-[[[1-[5-(trifluoromethyl)-2-pyridinyl]-1H-pyrazol-3-yl]oxy]methyl]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 175013-40-8 ZCAPLUS

CN Carbamic acid, methoxy[2-[[(1-pyrazinyl-1H-pyrazol-3-yl)oxy]methyl]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c} N \\ N \\ N \end{array} \begin{array}{c} N \\ N \\ M \in O - C - N \\ M \in O \end{array}$$

RN 175013-42-0 ZCAPLUS

CN Urea, N-methoxy-N'-methyl-N-[2-[[(1-pyrazinyl-1H-pyrazol-3-yl)oxy]methyl]phenyl]- (9CI) (CA INDEX NAME)

RN 175013-43-1 ZCAPLUS

CN Carbamic acid, methoxy[2-[[[4-nitro-1-[5-(trifluoromethyl)-2-pyridinyl]-1H-pyrazol-3-yl]oxy]methyl]phenyl]-, <math>methyl ester (9CI) (CA INDEX NAME)

L89 ANSWER 60 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1996:144848 ZCAPLUS Full-text

DOCUMENT NUMBER: 124:202243

TITLE: Preparation of methyl [alpha-(pyrazol-3-

yl)oxymethylene]phenylbutenoate agrochemical

fungicides and pesticides

INVENTOR(S): Oberdorf, Klaus; Koenig, Hartmann; Mueller, Bernd;

Kirstgen, Reinhard; Grammenos, Wassilios; Sauter,

Hubert; Lorenz, Gisela; Ammermann, Eberhard; Harries,

Volker

PATENT ASSIGNEE(S): Germany

SOURCE: PCT Int. Appl., 48 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

GΙ

PAT	PATENT NO.			KIND DATE			APPLICATION NO.					DATE			
WO	9529896			A1	19951109	,	 WO 19	995-EP15	54		1	99504	425	<	
	W: AU,	BG,	BR,	BY, CA	, CN, CZ,	FI,	HU,	JP, KR,	KΖ,	MX,	NO,	NΖ,	PL,		
	RU,	SG,	SK,	UA, US											
	RW: AT,	BE,	CH,	DE, DK	, ES, FR,	GB,	GR,	IE, IT,	LU,	MC,	NL,	PT,	SE		
IL	113414			A	20000813		IL 19	995-1134	14		1	99504	418	<	
CA	2189368			A1	19951109		CA 19	995-2189	368		1	99504	425	<	
AU	9524481			A	19951129		AU 19	995-2448	1		1	99504	425	<	
AU	682963			B2	19971023										
EP	758322			A1	19970219		EP 19	995-9186	03		1	99504	425	<	
EP	758322			B1	20010905										
	R: AT,	BE,	CH,	DE, DK	, ES, FR,	GB,	GR,	IE, IT,	LI,	NL,	PT,	SE			
CN	1150800			A	19970528	1	CN 19	995–1936	00		1	99504	425	<	
CN	1066137			В	20010523										
BR	9507602			A	19971007		BR 19	995-7602			1	99504	425	<	
JP	09512541			T	19971216	1	JP 19	995-5279	80		1	99504	425	<	
AT	205194			T	20010915		AT 19	995-9186	03		1	99504	425	<	
US	5707936			A	19980113		US 19	996-7323	00		1	99610	031	<	
PRIORITY	APPLN.	INFO.	. :				DE 19	994-4415	483		A 1	99405	503		
						,	WO 19	995-EP15	54	1	W 1	99504	425		
OTHER SC	OURCE(S):			MARPAT	124:2022	43									

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$$R^{2}$$

$$OCH_{2}$$

$$MeO - C - C - CHMe$$

The title compds. [I; n = 0-4; R1 = nitro, cyano, halogen, alkyl, haloalkyl, alkoxy; R2 = H, nitro, cyano, halogen, alkyl, haloalkyl, alkoxy, alkylthio, alkoxycarbonyl; R3 = (un)substituted alkyl, alkenyl, alkynyl; the dotted line represents an optional double bond], useful as agrochem. fungicides and pesticides, are prepared Thus, N-phenylpyrazolidin-3-one was condensed with Me α -(2- bromomethylphenyl)-2-butenoate, producing Me α -[2-(1-phenyl-4,5-dihydropyrazol-3-yloxymethyl)phenyl]-2-butenoate, m.p. 90-92°, which demonstrated agrochem. fungicidal activity against Plasmopara viticola.

Ι

174182-90-2P 174182-94-6P 174182-95-7P ΙT 174182-96-8P 174182-97-9P 174182-98-0P 174182-99-1P 174183-00-7P 174183-01-8P 174183-02-9F 174183-03-0P 174183-04-1P 174183-05-2P 174183-06-3P 174183-07-4P 174183-08-5P 174183-09-6P 174183-10-9P 174183-11-0P 174183-12-1P 174183-13-2P 174183-14-3P 174183-15-4P 174183-16-5P 174183-17-6P 174183-18-7P 174183-19-8P 174183-20-1P 174183-21-2P 174183-22-3P 174183-23-4P 174183-24-5P 174183-25-6P 174183-26-7P 174183-27-8P 174183-28-9P 174183-34-7P 174183-35-8P 174183-36-9P 174183-37-0P 174183-38-1P 174183-39-2P 174183-40-5P

RL: AGR (Agricultural use); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of Me [alpha-(pyrazol-3-yl))exymethylenelphenylbutenoate

(preparation of Me [alpha-(pyrazol-3-yl))oxymethylene]phenylbutenoate agrochem. fungicides and pesticides)

RN 174182-90-2 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(4-\text{chlorophenyl})-1H-\text{pyrazol}-3-\text{yl}] \text{oxy}] \text{methyl}] - \alpha-\text{ethylidene-, methyl ester (9CI)}$ (CA INDEX NAME)

RN 174182-94-6 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[(1-phenyl-1H-pyrazol-3-yl)oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 174182-95-7 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[[1-(4-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 174182-96-8 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(3-\text{chlorophenyl})-1H-\text{pyrazol}-3-\text{yl}] \text{oxy}] \text{methyl}] - \alpha-\text{ethylidene-, methyl ester (9CI)}$ (CA INDEX NAME)

RN 174182-97-9 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[[1-(4-fluorophenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 174182-98-0 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(2,4-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

RN 174182-99-1 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(2,4-dimethylphenyl)-1H-pyrazol-3-yl]oxy]methyl]-<math>\alpha$ -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-00-7 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(4-\text{chloro}-2-\text{methylphenyl})-1H-\text{pyrazol}-3-yl]oxy]\text{methyl}]-\alpha-\text{ethylidene-, methyl ester (9CI)}$ (CA INDEX NAME)

RN 174183-01-8 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(2-\text{chlorophenyl})-1H-\text{pyrazol}-3-\text{yl}] \text{oxy}] \text{methyl}]-\alpha-\text{ethylidene-, methyl ester (9CI)}$ (CA INDEX NAME)

RN 174183-02-9 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(3,5-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]-<math>\alpha$ -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-03-0 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(2,6-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-04-1 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[[1-(2-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-05-2 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[[1-(3-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-06-3 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[[1-(4-methoxyphenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-07-4 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(2,5-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]-<math>\alpha$ -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-08-5 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(3,4-\text{dichlorophenyl})-1\text{H-pyrazol}-3-y1] \text{oxy}] \text{methyl}]-\alpha-\text{ethylidene-, methyl ester (9CI)}$ (CA INDEX NAME)

RN 174183-09-6 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[[1-[3-(trifluoromethyl)phenyl]-

1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-10-9 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[[1-[5-(trifluoromethyl)-2-pyridinyl]-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-11-0 ZCAPLUS

CN Benzeneacetic acid, $2-[[[4-\text{chloro}-1-(4-\text{methylphenyl})-1H-\text{pyrazol}-3-yl] \text{oxy}] \text{methyl}]-\alpha-\text{ethylidene-, methyl ester (9CI)}$ (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 174183-12-1 ZCAPLUS

CN Benzeneacetic acid, $2-[[[4-chloro-1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]-<math>\alpha$ -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & &$$

RN 174183-13-2 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[[1-(3-methoxyphenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-14-3 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(2,2-difluoro-2,3-dihydro-1H-inden-5-yl)-1H-pyrazol-3-yl]oxy]methyl]- α -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-15-4 ZCAPLUS

CN 1H-Pyrazole-4-carboxylic acid, 1-(4-chlorophenyl)-3-[[2-[1-(methoxycarbonyl)-1-propenyl]phenyl]methoxy]-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-16-5 ZCAPLUS

CN 1H-Pyrazole-4-carboxylic acid, 1-(2,4-dichlorophenyl)-3-[[2-[1-(methoxycarbonyl)-1-propenyl]phenyl]methoxy]-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-17-6 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[[1-[4-(trifluoromethyl)phenyl]-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-18-7 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[[1-(2-pyridinyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-19-8 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[(5-methyl-1-phenyl-1H-pyrazol-3-yl)oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-20-1 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(2,4-\text{dichlorophenyl})-5-(\text{trifluoromethyl})-1H-pyrazol-3-yl]oxy]methyl]-<math>\alpha$ -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-21-2 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(2,4-\text{dichlorophenyl})-4-\text{nitro-1H-pyrazol-3-yl]oxy]methyl]-\alpha-ethylidene-, methyl ester (9CI) (CA INDEX NAME)$

RN 174183-22-3 ZCAPLUS

CN Benzeneacetic acid, $2-[[[4-chloro-1-(2,4-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]-<math>\alpha$ -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-23-4 ZCAPLUS

CN 1H-Pyrazole-4-carboxylic acid, 3-[[2-[1-(methoxycarbonyl)-1-propenyl]phenyl]methoxy]-1-phenyl-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-24-5 ZCAPLUS

CN Benzeneacetic acid, 2-[[[4-chloro-1-[5-(trifluoromethyl)-2-pyridinyl]-1H-pyrazol-3-yl]oxy]methyl]- α -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-25-6 ZCAPLUS

CN Benzeneacetic acid, 2-chloro-6-[[[1-(2,4-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-26-7 ZCAPLUS

CN Benzeneacetic acid, 2-chloro-6-[[[1-(3,5-dichlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]- α -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-27-8 ZCAPLUS

CN Benzeneacetic acid, 2-chloro- α -ethylidene-6-[[[1-(4-methoxyphenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-28-9 ZCAPLUS

CN Benzeneacetic acid, 2-chloro- α -ethylidene-6-[[[1-[5-(trifluoromethyl)-2-pyridinyl]-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

$$F_3C$$
N
O-CH2
C1
C-CH-Me
C-OMe

RN 174183-34-7 ZCAPLUS

CN Benzeneacetic acid, $2-[[[1-(5-\text{chloro}-2-\text{pyridinyl})-1\text{H-pyrazol}-3-yl] \text{oxy}] \text{methyl}]-\alpha-\text{ethylidene-, methyl ester (9CI)}$ (CA INDEX NAME)

RN 174183-35-8 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(6-chloro-3-pyridazinyl)-1H-pyrazol-3-yl]oxy]methyl]- α -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-36-9 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[(1-pyrazinyl-1H-pyrazol-3-yl)oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-37-0 ZCAPLUS

CN Benzeneacetic acid, 2-[[(4-chloro-1-phenyl-1H-pyrazol-3-yl)oxy]methyl]- α -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-38-1 ZCAPLUS

CN Benzeneacetic acid, 2-[[(4-chloro-1-pyrazinyl-1H-pyrazol-3-yl)oxy]methyl]- α -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

RN 174183-39-2 ZCAPLUS

CN Benzeneacetic acid, $2-[[[4-bromo-1-(4-chlorophenyl)-1H-pyrazol-3-yl]oxy]methyl]-<math>\alpha$ -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-40-5 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[[4-nitro-1-[5-(trifluoromethyl)-2-pyridinyl]-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

IT 174182-93-5

RL: RCT (Reactant); RACT (Reactant or reagent) (preparation of Me [alpha-(pyrazol-3-yl)oxymethylene]phenylbutenoate agrochem. fungicides and pesticides)

RN 174182-93-5 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[[1-(6-fluoro-2-pyridinyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

IT 174183-42-7P 174183-43-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of Me [alpha-(pyrazol-3-yl)oxymethylene]phenylbutenoate agrochem. fungicides and pesticides)

RN 174183-42-7 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(2-chloro-4-fluorophenyl)-1H-pyrazol-3-

yl]oxy]methyl]- α -ethylidene-, methyl ester (9CI) (CA INDEX NAME)

RN 174183-43-8 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[[1-(4-fluoro-2-methylphenyl)-1H-pyrazol-3-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

L89 ANSWER 61 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1995:995024 ZCAPLUS Full-text

DOCUMENT NUMBER: 124:117306

TITLE: Preparation of pyrazolyloxymethylphenylpropenoic ester

derivatives as agrochemical fungicides

INVENTOR(S): Hwang, Ku-Jun; Kim, Sung Soo; Kim, Byung Sup

PATENT ASSIGNEE(S): Korea Research Institute of Chemical Technology, S.

Korea

SOURCE: PCT Int. Appl., 28 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
		1005001		10050014
WO 9525095	A1	19950921	WO 1995-KR20	19950314 <
W: AU, BR, C	A, JP, US	5		
RW: AT, BE, C	H, DE, DH	K, ES, FR,	GB, GR, IE, IT, LU, MC	, NL, PT, SE
KR 9706238	B1	19970425	KR 1994-5088	19940315 <
AU 9519617	A	19951003	AU 1995-19617	19950314 <
AU 692847	B2	19980618		
EP 750613	A1	19970102	EP 1995-912491	19950314 <
R: DE, ES, F	R, GB, II	- -		
JP 09503525	T	19970408	JP 1995-523958	19950314 <
JP 3111320	B2	20001120		
US 5776965	A	19980707	US 1996-702634	19961101 <

PRIORITY APPLN. INFO.: KR 1994-5088 A 19940315

WO 1995-KR20 W 19950314

OTHER SOURCE(S): MARPAT 124:117306

GT

$$R^{1}$$
 $CO2Me$
 R^{5}
 R^{4}
 R^{5}
 R^{6}
 R^{6}
 R^{1}
 R^{6}
 R^{7}
 R^{1}
 R^{1}
 R^{2}
 R^{1}
 R^{2}
 R^{1}
 R^{2}
 R^{2}
 R^{2}
 R^{2}
 R^{3}
 R^{4}
 R^{5}
 R^{5}
 R^{6}
 R^{6}
 R^{6}
 R^{7}

AΒ The title compds. I [R1 represents hydrogen, halogen, nitro, an alkyl group having 1 to 6 carbon atoms, or an alkoxy group having 1 to 6 carbon atoms; R2 represents an alkoxy group having 1 to 6 carbon atoms, a haloalkoxy group having 1 to 6 carbon atoms, or an alkylthio group having 1 to 6 carbon atoms; R3 represents an alkyl group having 1 to 6 carbon atoms, an allyl group, a benzyl group, a Ph group, or a substituted Ph group by substituent selected from the group consisting of an alkyl group having 1 to 6 carbon atoms, an alkoxy group having 1 to 6 carbon atoms, nitro and halogen; R4 represents hydrogen, halogen, an alkyl group having 1 to 6 carbon atoms, a Ph group, etc.; R5 represents hydrogen, halogen, a haloalkyl group, etc.; and X represents carbon or nitrogen] are claimed. The title compound trans-II was prepared from Me 2-(2-bromomethylphenyl)-3- methoxypropenoate and 1-methyl-3trifluoromethyl-4-hydroxypyrazole. Trans-II showed EC50 of 250 ppm against rice blast. Five other compds. of this invention showed EC50 values of $<2\ \text{ppm}$ to 10 ppm against rice blast, wheat leaf rust, etc.

172834-81-0P 172834-82-1P 172834-83-2P ΙT

172834-84-3P 172834-85-4P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of pyrazolyloxymethylphenylpropenoic ester derivs. as

agrochem.

fungicides)

172834-81-0 ZCAPLUS RN

CN Benzeneacetic acid, α -(methoxymethylene)-2-[[[1-phenyl-3-(trifluoromethyl)-1H-pyrazol-4-yl]oxy]methyl]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

$$F_3C$$

$$Ph$$

$$MeO$$

$$E$$

$$OMe$$

RN 172834-82-1 ZCAPLUS

CN Benzeneacetic acid, α -(methoxyimino)-2-[[[1-phenyl-3-(trifluoromethyl)-1H-pyrazol-4-yl]oxy]methyl]-, methyl ester, (Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 172834-83-2 ZCAPLUS

CN Benzeneacetic acid, α -(methoxyimino)-2-[[[1-phenyl-3-(trifluoromethyl)-1H-pyrazol-4-yl]oxy]methyl]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

$$F_3 \subset \bigvee^{\text{Ph}} \bigvee^{\text{MeO}} \bigvee^{\text{O}} \bigvee^{\text{E}} \bigvee^{\text{OMe}}$$

RN 172834-84-3 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[[(3-methyl-1-phenyl-1H-pyrazol-4-yl)oxy]methyl]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 172834-85-4 ZCAPLUS

CN Benzeneacetic acid, α -[(methylthio)methylene]-2-[[[1-phenyl-3-

(trifluoromethyl)-1H-pyrazol-4-yl]oxy]methyl]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

$$\mathsf{F}_3\mathsf{C} \qquad \mathsf{MeO} \qquad \mathsf{E} \qquad \mathsf{SMe}$$

IT 172834-86-5

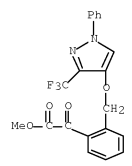
RL: RCT (Reactant); RACT (Reactant or reagent)

(preparation of pyrazolyloxymethylphenylpropenoic ester derivs. as agrochem.

fungicides)

RN 172834-86-5 ZCAPLUS

CN Benzeneacetic acid, α -oxo-2-[[[1-phenyl-3-(trifluoromethyl)-1H-pyrazol-4-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)



L89 ANSWER 62 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1995:731727 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 123:112056

TITLE: 5-Arylisoxazol-4-yl-substituted 2-amino carboxylic

acid compounds

INVENTOR(S): Moltzen, Lenz Sibylle; Falch, Erik; Boegesoe, Klaus

Peter; Krogsgaard-Larsen, Povl

PATENT ASSIGNEE(S): H. Lundbeck A/S, Den. SOURCE: PCT Int. Appl., 54 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9512587	A1	19950511	WO 1994-DK411	19941102 <

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W: AM, AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, ES, FI, GB,
             GE, HU, JP, KE, KG, KP, KR, KZ, LK, LT, LU, LV, MD, MG, MN, MW,
             NL, NO, NZ, PL, PT, RO, RU, SD, SE, SI, SK, TJ, TT, UA, US, UZ, VN
         RW: KE, MW, SD, SZ, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU,
             MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN,
     CA 2175685
                                19950511
                                            CA 1994-2175685
                                                                    19941102 <--
                          Α1
     AU 9480579
                                             AU 1994-80579
                                                                    19941102 <--
                                19950523
                          Α
     AU 680062
                          В2
                                19970717
                                             ZA 1994-8631
     ZA 9408631
                          Α
                                19950710
                                                                    19941102 <--
     EP 726896
                                            EP 1994-931523
                                19960821
                                                                    19941102 <--
                          Α1
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE
     CN 1136810
                                            CN 1994-194388
                          Α
                                19961127
                                                                    19941102 <--
     CN 1056837
                                20000927
                          R
     HU 74692
                                            HU 1996-1167
                                                                    19941102 <--
                          Α2
                                19970128
     JP 09504531
                          Τ
                                19970506
                                             JP 1994-512970
                                                                    19941102 <--
     RU 2138488
                          C1
                                19990927
                                             RU 1996-112168
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                                20000419
     EP 994107
                          Α1
                                            EP 1999-125828
                                                                    19941102 <--
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT
     FI 9601872
                                19960503
                                             FI 1996-1872
                                                                    19960502 <--
                          Α
     NO 9601783
                                19960625
                                             NO 1996-1783
                                                                    19960502 <--
                          Α
PRIORITY APPLN. INFO.:
                                             DK 1993-1243
                                                                 A 19931103
                                             EP 1994-931523
                                                                 A3 19941102
                                             WO 1994-DK411
                                                                 W 19941102
```

OTHER SOURCE(S): MARPAT 123:112056

GΙ

AΒ 2-Aminocarboxylic acid compds. substituted with 5-arylisoxazol-4-yl or 5arylisothiazol-4-yl groups are claimed, specifically compds. I [A = bond or]spacer; B = group CH(NR'R'')CO2H where R' and R'' = H or C1-6 alkyl, or B = cyclobutenedione group Q wherein R2, R3 and R4 = various substituents; or R3R4 or R2R4 form ring; E = O, S, CO2, (CH2)nCO2, O(CH2)nCO2, or S(CH2)nCO2 wherein n = 1-6, 5-tetrazolyl, 5-tetrazolylalkyl, 3-hydroxyisoxazolyl, or 3hydroxyisoxazolylalkyl; D = O or S; R1 = (un)substituted aryl or heteroaryl; certain racemic forms excluded]. I are excitatory amino acid receptor ligands useful in the treatment of cerebral ischemia, Huntington's disease, epileptic disorders, Parkinson's disease, Alzheimer's disease, schizophrenia, pain, depression and anxiety. For example, cyanation of 2-bromothiophene with CuCN in refluxing NMP gave 63% 2-thiophenecarbonitrile, which reacted with MeCHBrCO2Et and Zn in the presence of CuBr2 to give 72% Et 2-methyl-3-(2thienyl)-3-oxopropionate. This was cyclized with NH2OH to give 55% isoxazole derivative II (G3 = OH, G4 = Me), which underwent O-ethylation with EtBr and

K2CO3 (51%) and benzylic bromination with NBS (100%) to give II (G3 = OEt, G4 = CH2Br). The latter was used to alkylate AcNHCH(CO2Et)2 (68%), and the resulting malonate diester was saponified, decarboxylated, deacetylated, and deethylated in refluxing 48% HBr, to give 30% title compound (±)-III. In the cortical wedge model in rats, this compound showed an AMPA agonist profile, with an EC50 of 5.8 μM . A variety of addnl. I were similarly prepared and tested by this and other binding assays; they showed activity as agonists or antagonists at NMDA and/or AMPA receptors.

IT 166180-57-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate; preparation of arylisoxazolyl amino carboxylic acids as AMPA/NMDA receptor ligands)

RN 166180-57-0 ZCAPLUS

CN Benzoic acid, 2-[[[[3-(carboxymethoxy)-5-(2-thienyl)-4-isoxazolyl]methyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

L89 ANSWER 63 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1995:229475 ZCAPLUS Full-text

DOCUMENT NUMBER: 122:239694

TITLE: Pesticidal 1-aryl-5-(substituted

alkylideneimino)pyrazoles

INVENTOR(S): Huang, Jamin; Ayad, Hafez M.; Timmons, Philip R.

PATENT ASSIGNEE(S): Rhone-Poulenc AG Co., USA

SOURCE: U.S., 24 pp. Cont.-in-part of U.S. Ser. No. 790,449,

abandoned.
CODEN: USXXAM

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PA	TENT NO.	KIND		D DATE	APPLICATION NO.	DATE
US	5360910		 A	19941101	US 1992-842431	19920304 <
US	5236938		А	19930817	US 1991-693580	19910430 <
CA	2067282		A1	19921031	CA 1992-2067282	19920427 <
AU	9215192		А	19921105	AU 1992-15192	19920427 <
AU	655014		В2	19941201		
IL	101702		А	19960331	IL 1992-101702	19920427 <
ИО	9201639		А	19921102	NO 1992-1639	19920428 <
ИО	303631		В1	19980810		
EP	511845		A1	19921104	EP 1992-303857	19920429 <
ΕP	511845		В1	20011031		
	R: AT,	BE, CH	, DE,	DK, ES, FR,	GB, GR, IT, LI, LU,	NL, PT, SE
HU	61529		A2	19930128	HU 1992-1416	19920429 <
HU	213630		В	19970828		

69737	В1	19960830	PL	1992-294383		19920429	<
2088576	C1	19970827	RU	1992-5011630		19920429	<
207904	T	20011115	ΑT	1992-303857		19920429	<
2165353	Т3	20020316	ES	1992-303857		19920429	<
511845	T	20020429	PΤ	1992-303857		19920429	<
.066265	A	19921118	CN	1992-103156		19920430	<
.053659	В	20000621					
201735	A	19921124	BR	1992-1735		19920430	<
203175	A	19930127	ZA	1992-3175		19920430	<
)5148240	A	19930615	JΡ	1992-111958		19920430	<
3248943	B2	20020121					
07407	В1	19931130	RO	1992-598		19920430	<
279252	В6	19980805	SK	1992-1337		19920430	<
286232	В6	20000216	CZ	1992-1337		19920430	<
APPLN. INFO.:			US	1991-693580	Α2	19910430	
			US	1991-790449	В2	19911112	
			US	1992-842431	Α	19920304	
			CS	1992-1337	A	19920430	
	088576 07904 165353 11845 066265 053659 201735 203175 5148240 248943 07407 79252 86232	088576 C1 07904 T 165353 T3 11845 T 066265 A 053659 B 201735 A 203175 A 5148240 A 248943 B2 07407 B1 79252 B6 86232 B6	088576 C1 19970827 07904 T 20011115 165353 T3 20020316 11845 T 20020429 066265 A 19921118 053659 B 20000621 201735 A 19921124 203175 A 19930127 5148240 A 19930615 248943 B2 20020121 07407 B1 19931130 79252 B6 19980805 86232 B6 20000216	088576 C1 19970827 RU 07904 T 20011115 AT 165353 T3 20020316 ES 11845 T 20020429 PT 066265 A 19921118 CN 053659 B 20000621 201735 A 19921124 BR 203175 A 19930127 ZA 5148240 A 19930615 JP 248943 B2 20020121 07407 B1 19931130 RO 79252 B6 19980805 SK 86232 B6 20000216 CZ APPLN. INFO.: US	088576 C1 19970827 RU 1992-5011630 07904 T 20011115 AT 1992-303857 165353 T3 20020316 ES 1992-303857 11845 T 20020429 PT 1992-303857 066265 A 19921118 CN 1992-103156 053659 B 20000621 201735 A 19921124 BR 1992-1735 203175 A 19930127 ZA 1992-3175 5148240 A 19930615 JP 1992-111958 248943 B2 20020121 07407 B1 19931130 RO 1992-598 79252 B6 19980805 SK 1992-1337 86232 B6 20000216 CZ 1992-1337	088576 C1 19970827 RU 1992-5011630 07904 T 20011115 AT 1992-303857 165353 T3 20020316 ES 1992-303857 11845 T 20020429 PT 1992-303857 066265 A 19921118 CN 1992-103156 053659 B 20000621 201735 A 19921124 BR 1992-1735 203175 A 19930127 ZA 1992-3175 5148240 A 19930615 JP 1992-111958 248943 B2 20020121 07407 B1 19931130 RO 1992-598 79252 B6 19980805 SK 1992-1337 79252 B6 19980805 SK 1992-1337 86232 B6 20000216 CZ 1992-1337 APPLN. INFO.: US 1991-693580 A2 US 1991-790449 B2 US 1992-842431 A	088576

OTHER SOURCE(S):

MARPAT 122:239694

GΙ

AΒ The invention describes novel 1-aryl-5-(substituted alkylideneimino)pyrazole of formula (I) wherein typically preferred substituents are: R1 is cyano, nitro, or halogen; R2 is R9S(O)n in which n is 0, 1 or 2 and R9 is alkyl, preferably Me which is substituted by halogen atoms which are the same or different up to full substitution of the alkyl moiety; R3 is hydrogen or alkyl; R4 is Ph or heteroaryl, optionally substituted by one or more hydroxy, halogen, alkoxy, alkylthio, cyano or alkyl or combinations thereof; preferably R4 is Ph, which is at least substituted by 3-hydroxy or 4-hydroxy; R5 is hydrogen, alkyl or halogen; R6 and R8 are hydrogen; R7 is halogen, alkyl, haloalkyl or haloalkoxy; and X is a nitrogen atom or CR14 in which R14 is hydrogen, halogen, cyano, alkyl, alkylthio or alkoxy. The invention further describes processes to make the compds., compns. of the compds., and methods of use of the compds. for the control of arthropods (mites, aphids or insects), nematodes, helminths, or protozoa. Pesticidal activity of I compds. providing 70-100% pest mortality was evaluated against buckthorn aphid, cotton aphid, southern armyworm, Mexican bean beetle, housefly, tobacco budworm, southern corn rootworm, western corn rootworm. ΙT

162368-35-6P 162368-36-7P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(pesticidal 1-aryl-5-(substituted alkylideneimino)pyrazoles)

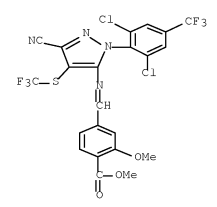
162368-35-6 ZCAPLUS RN

CN Benzoic acid, 4-[[[3-cyano-1-[2,6-dichloro-4-(trifluoromethyl)phenyl]-4-

[(trifluoromethyl)thio]-1H-pyrazol-5-yl]imino]methyl]-, methyl ester (9CI)
 (CA INDEX NAME)

RN 162368-36-7 ZCAPLUS

CN Benzoic acid, 4-[[[3-cyano-1-[2,6-dichloro-4-(trifluoromethyl)phenyl]-4[(trifluoromethyl)thio]-1H-pyrazol-5-yl]imino]methyl]-2-methoxy-, methyl
ester (9CI) (CA INDEX NAME)



L89 ANSWER 64 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1995:220182 ZCAPLUS Full-text

DOCUMENT NUMBER: 122:9667

TITLE: Preparation of α -(2-ethenylphenyl)acrylates as

pesticides

INVENTOR(S): Kirstgen, Reinhard Dr; Theobald, Hans Dr; Oberdorf,

Klaus Dr; Doetzer, Reinhard Dr; Klintz, Ralf Dr;

Schaefer, Bernd Dr; Harries, Volker Dr; Kardorff, Uwe

Dr; Lorenz, Gisela Dr; Ammermann, Eberhard Dr

PATENT ASSIGNEE(S): BASF A.-G., Germany SOURCE: Ger. Offen., 133 pp.

CODEN: GWXXBX

CODEN: GWXXB

DOCUMENT TYPE: Patent
LANGUAGE: German
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:	TENT	NO.			KIN	D	DATE		APPLICATION NO.						DATE			
CA	4238 2149 9411 W:	238 334 AU,	BB,	BG,	A1 A1 BR,	BY,	1994 1994 , CA,	0526 0526 CZ,	FI,	CA 1 WO 1 HU,	993- 993- JP,	2149 EP30 KP,	238 67 KR,	KΖ,	1 1	9931	102 102	<
	R₩:	AT,	BE,	CH,	DE,	DK	, RO, , ES, , CM,	FR,	GB,	GR,	IE,	ΙT,	LU,	MC,		PT,	SE,	,
AU	9454	634			Α		1994	0608		AU 1	994-	5463	4		1	9931	102	<
AU	6715	04			В2		1996	0829										
EP	6688	52			A1		1995	0830		EP 1	994-	9000	87		1	9931	102	<
EP	6688	52			В1		1998	0225										
	R:	ΑT,																
	7315	-			A2		1996											
	0850												73					
	1634				T		1998									9931		
	2114				Т3		1998						87			9931		
IL	1075	20			Α		1998	1206		IL 1	993-	1075	20		1	9931	105	<
ZA	9308	414			Α		1995	0511		ZA 1	993-	8414			1	9931	111	<
	1098				Α		1995	0215		CN 1	993-	1213	27		1	9931	112	<
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US	5633	268			Α		1997	0527		US 1	995-	4335	15		1	9950	512	<
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										WO 1	993-	EP30	67		W 1	9931	102	
OTHER SO	OURCE	(S):			MAR:	PAT	122:	9667										

AB R2R3C:CHZC(:X)COYMe [R2 = NO2, cyano, halo, alkoxy, alkanoylamino, alkoxycarbonylamino, NHCO2CH2Ph; R3 = halo, (hetero)aryl, C(:Z1)TR4, C(:Z2)R5; R4 = H, alk(en)yl, aryl, etc.; R5 = H, cyano, halo, alkyl, alkoxy, aryl, etc.; T = O, S, NH, etc.; X = CHOMe, CHMe, NOMe; Y = O or NH; Z = (un)substituted 1,2-C6H4; Z1 = O, S, (alkyl)imino, etc.; Z2 = O, (alkyl)imino, hydrazono, etc.] were prepared as agrochem. fungicides, insecticides, acaricides, and nematocides. Thus, (E)-2-(OHC)C6H4C(:CHOMe)CO2Me was condensed with (EtO)2P(O)CHClCO2Me to give title compound I (R = OMe) which was converted in 3 steps to I (R = SCMe3). The latter gave $\geq 80\%$ control of Aphis fabae at 200ppm.

IT 159375-84-5P 159375-85-6P 159375-86-7P 159375-87-8P 159375-88-9P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as pesticide)

RN 159375-84-5 ZCAPLUS

CN Benzeneacetic acid, 4-chloro-2-[2-chloro-2-[3-(4-chlorophenyl)-5-

isoxazolyl]ethenyl]- α -(methoxymethylene)-, methyl ester, (Z,E)-(9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 159375-85-6 ZCAPLUS

CN Benzeneacetic acid, 4-chloro-2-[2-chloro-2-[3-(4-chlorophenyl)-5-isoxazolyl]ethenyl]- α -(methoxymethylene)-, methyl ester, (E,E)-(9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 159375-86-7 ZCAPLUS

CN Benzeneacetic acid, 2-[2-chloro-2-[3-(4-chlorophenyl)-5-isoxazolyl]ethenyl]- α -(methoxymethylene)-, methyl ester, (Z,E)-(9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 159375-87-8 ZCAPLUS

CN Benzeneacetic acid, 2-[2-chloro-2-[3-(4-chlorophenyl)-5-isoxazolyl]ethenyl]-5-(1,1-dimethylethyl)- α -(methoxymethylene)-, methyl ester, (Z,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 159375-88-9 ZCAPLUS

CN Benzeneacetic acid, 2-[2-chloro-2-[4-chloro-3-(4-chlorophenyl)-5-isoxazolyl]ethenyl]- α -(methoxymethylene)-, methyl ester, (Z,E)-(9CI) (CA INDEX NAME)

Double bond geometry as shown.

L89 ANSWER 65 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1994:655793 ZCAPLUS Full-text

ACCESSION NUMBER: 1994:000/90 ZCAPLOS EUII-

DOCUMENT NUMBER: 121:255793

TITLE: Preparation of ortho-substituted N-methyl- α -

 $(\verb"methoxyimino") \verb"benzeneacetam" ides as fungicides or$

insecticides

INVENTOR(S): Kirstgen, Reinhard; Grammenos, Wassilios; Bayer,

Herbert; Doetzer, Reinhard; Koenig, Hartmann;
Oberdorf, Klaus; Sauter, Hubert; Wingert, Horst;

Lorenz, Gisela; et al.

PATENT ASSIGNEE(S): BASF A.-G., Germany SOURCE: Ger. Offen., 56 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 4305502	A1	19940825	DE 1993-4305502	19930223 <
IL 108462	A	19981030	IL 1994-108462	19940128 <
CA 2155571	A1	19940901	CA 1994-2155571	19940212 <
WO 9419331	A1	19940901	WO 1994-EP408	19940212 <
W: AU, BB, BG,	BR, BY	, CA, CN, CZ	I, FI, GE, HU, JP, KP,	KR, KZ, LK,

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LV, MG, MN, MW, NO, NZ, PL, RO, RU, SD, SK, UA, US, UZ, VN
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
             BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
     AU 9461091
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                                19940914
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     AU 682339
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         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, NL, PT, SE
                                            BR 1994-5937
                                                                    19940212 <--
     BR 9405937
                          Α
                                19960206
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                                            CN 1994-191272
                                                                    19940212 <--
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                                19960306
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                                                                    19940212 <--
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                                            HU 1995-2454
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     ES 2102206
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                                19970716
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     RU 2130924
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                                19990527
                                            RU 1995-122817
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     PL 179860
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                                20001130
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    AT 249445
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PRIORITY APPLN. INFO.:
                                             DE 1993-4305502
                                                                 A 19930223
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                                            WO 1994-EP408
                                                                 W 19940212
                                                                 A3 19950821
                                             US 1995-505288
OTHER SOURCE(S):
                        MARPAT 121:255793
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GΙ

AB The title compds., ortho-substituted N-methyl- α - (methoxyimino) benzeneacetamides I (R1 = nitro, cyano, halo, alkyl, etc.; R2 = H, alkyl, etc.; X = oxygen, sulfur; Y = heteroarom. ring; n = integer) were disclosed as fungicides, insecticides, acaricides and nematocides. An example compound, (E)- α -(methoxyimino)-2-[2-[1-(4-chloro-2-methylphenyl)-1H-pyrazol-4-yl]oxymethyl]-N-methylbenzeneacetamide (II) was prepared Biol. test data for I were not shown.

IT 158668-39-4P 158668-47-4P 158668-48-5P 158668-49-6P 158668-50-9P 158668-51-0P 158668-52-1P 158668-53-2P 158668-54-3P

158668-55-4P

RL: AGR (Agricultural use); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of α -(methoxyimino)benzeneacetamides as fungicides insecticides)

RN 158668-39-4 ZCAPLUS

CN Benzeneacetamide, 2-[[[5-(4-chlorophenyl)-3-isoxazolyl]oxy]methyl]- α -(methoxyimino)-N-methyl- (9CI) (CA INDEX NAME)

RN 158668-47-4 ZCAPLUS

CN Benzeneacetamide, α -(methoxyimino)-N-methyl-2-[[(1-phenyl-1H-pyrazol-4-yl)oxy]methyl]- (9CI) (CA INDEX NAME)

RN 158668-48-5 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(4-fluorophenyl)-1H-pyrazol-4-yl]oxy]methyl]- α -(methoxyimino)-N-methyl- (9CI) (CA INDEX NAME)

RN 158668-49-6 ZCAPLUS

CN Benzeneacetamide, α -(methoxyimino)-N-methyl-2-[[[1-[4-(trifluoromethyl)phenyl]-1H-pyrazol-4-yl]oxy]methyl]- (9CI) (CA INDEX NAME)

RN 158668-50-9 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(4-cyanophenyl)-1H-pyrazol-4-yl]oxy]methyl]- α -(methoxyimino)-N-methyl- (9CI) (CA INDEX NAME)

RN 158668-51-0 ZCAPLUS

CN Benzeneacetamide, α -(methoxyimino)-N-methyl-2-[[[1-(4-methylphenyl)-1H-pyrazol-4-yl]oxy]methyl]- (9CI) (CA INDEX NAME)

RN 158668-52-1 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(4-chlorophenyl)-1H-pyrazol-4-yl]oxy]methyl]- α -(methoxyimino)-N-methyl- (9CI) (CA INDEX NAME)

RN 158668-53-2 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(2,4-dichlorophenyl)-1H-pyrazol-4-yl]oxy]methyl]- α -(methoxyimino)-N-methyl- (9CI) (CA INDEX NAME)

RN 158668-54-3 ZCAPLUS

CN Benzeneacetamide, 2-[[[1-(4-chloro-2-methylphenyl)-1H-pyrazol-4-yl]oxy]methyl]- α -(methoxyimino)-N-methyl-, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

$$\begin{array}{c} \text{MeO} \\ \text{N} \\ \text{E} \\ \text{NHMe} \\ \text{N} \\ \text{Me} \\ \text{N} \\ \text{Me} \\ \text{N} \\ \text{Me} \\ \text{N} \\ \text$$

RN 158668-55-4 ZCAPLUS

CN Benzeneacetamide, 2-[[[5-(4-chlorophenyl)-3-isoxazolyl]oxy]methyl]- α -(methoxyimino)-N-methyl-, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

IT 158668-57-6P

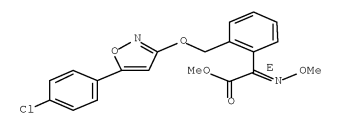
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of α -(methoxyimino)benzeneacetamides as fungicides insecticides)

RN 158668-57-6 ZCAPLUS

CN Benzeneacetic acid, $2-[[[5-(4-\text{chlorophenyl})-3-\text{isoxazolyl}] \text{oxy}] \text{methyl}] - \alpha-(\text{methoxyimino})-, \text{methyl ester, (E)- (9CI)}$ (CA INDEX NAME)

Double bond geometry as shown.



L89 ANSWER 66 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1994:604982 ZCAPLUS Full-text

DOCUMENT NUMBER: 121:204982

TITLE: Acetylenic derivatives and their use as

plant-protective agents

INVENTOR(S): Wingert, Horst; Hellendahl, Beate; Kirstgen, Reinhard;

Sauter, Hubert; Ammermann, Eberhard; Lorenz, Gisela

PATENT ASSIGNEE(S): BASF A.-G., Germany SOURCE: Eur. Pat. Appl., 70 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAI	ENT	NO.			KINI	O	DATE		AP	PLICA	NOITA	NO.		DATE		
	5829				A1		1994		EP	1993	3-1123	327		1993073	31	<
EP	5829 R:		BE,	CH,	B1 DE,		1996 ES,		GB, G	R, II	E, IT,	, LI,	NL,	PT, SE		
CA	2101	664			A1		1994	0212	CA	1993	3-2101	1664		1993073	30	<
US	5449	809			Α		1995	0912	US	1993	3-9969	9.3		1993073	30	<

	718292 718292		A1 B1	19960626 19980422	EP 1996-101050 19	930731 <
ш	R: AT,	BE. CH.			GB, GR, IE, IT, LI, NL, PT,	SE
АТ	143657	DL, 011,	T T	19961015		930731 <
ES	2093335		T3	19961216	ES 1993-112327 19	930731 <
AT	165352		T	19980515	AT 1996-101050 19	930731 <
AU	9344517		A	19940217	AU 1993-44517 19	930810 <
AU	663208		B2	19950928		
JP	06239824		A	19940830	JP 1993-198510 19	930810 <
ZA	9305787		A	19950210	ZA 1993-5787 19	930810 <
HU	68742		A2	19950728	HU 1993-2315 19	930810 <
US	5686474		A	19971111	US 1995-443460 19	950518 <
PRIORITY	APPLN.	INFO.:			DE 1992-4226557 A 19	920811
					DE 1992-4239874 A 19	921127
					US 1993-99693 A3 19	930730
					EP 1993-112327 A3 19	930731
OTHER SC GI	OURCE(S):		MARPAT	121:20498	2	

$$\begin{array}{c} U & V \\ V & W \\ C = CR \end{array}$$

RN

AB Title acetylene derivs. of general formula I, wherein U, V and W can be the same or different and are selected from H, halogen, nitro, cyano, or alkyl or alkoxy of 1 to 4 carbon atoms, A = alkylidene, alkylthio- or alkoxymethylidene or alkoxymino of 1 to 4 carbon atoms, B = OH, alkoxy and alkylamino of 1 to 4 carbon atoms, R = e.g., H, halogen, CF3, alkyl, heteroarylthiomethyl, etc. were prepared and tested for fungicidal activity.

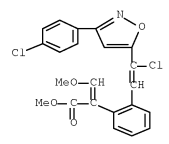
IT 158036-25-OP 158036-30-7P

158036-25-0P 158036-30-7P
RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
 (preparation and reactivity of, as plant-protective fungicidal agents)
158036-25-0 ZCAPLUS

CN Benzeneacetic acid, 2-[[3-(4-chlorophenyl)-5-isoxazolyl]ethynyl]- α - (methoxymethylene)-, methyl ester (9CI) (CA INDEX NAME)

RN 158036-30-7 ZCAPLUS

CN Benzeneacetic acid, 2-[2-chloro-2-[3-(4-chlorophenyl)-5-isoxazolyl]ethenyl]- α -(methoxymethylene)-, methyl ester (9CI) (CA INDEX NAME)



L89 ANSWER 67 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1994:469414 ZCAPLUS Full-text

DOCUMENT NUMBER: 121:69414

TITLE: Silver halide photographic material containing

antiirradiation dye and polymer latex to improve

quality of printed characters

INVENTOR(S): Morihara, Hideaki; Yoshida, Kazuhiro; Arai, Takeo

PATENT ASSIGNEE(S): Konishiroku Photo Ind, Japan SOURCE: Jpn. Kokai Tokkyo Koho, 22 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 06035097	A	19940210	JP 1992-195444	19920722 <
PRIORITY APPLN. INFO.:			JP 1992-195444	19920722

- AB The claimed photog. material having ≥1 light-sensitive layer and ≥1 light-insensitive hydrophilic colloid layer on a support is characterized by (1) that the emulsion layer and the colloid layer contain a polymer latex stabilized by gelatin and (2) that the emulsion layer and/or hydrophilic colloid layer contains a water-soluble dye having the absorption peak at 400-500 nm. It provides a printed characters with an excellent sharpness with low background d., and remains little residual dye stain in the processed materials.
- IT 156245-66-8

RL: TEM (Technical or engineered material use); USES (Uses) (photog. material containing, antiirradn. dye)

RN 156245-66-8 ZCAPLUS

CN 1H-Pyrazole-1-acetic acid, 3-ethyl-4-[[3-ethyl-5-hydroxy-1-(4-sulfophenyl)-1H-pyrazol-4-yl]methylene]-4,5-dihydro-5-oxo-, disodium salt (9CI) (CA INDEX NAME)

●2 Na

L89 ANSWER 68 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1994:408893 ZCAPLUS Full-text

DOCUMENT NUMBER: 121:8893

TITLE: Phenyl-substituted acrylate ester agrochemical

fungicides

INVENTOR(S):
Mueller, Bernd; Roehl, Franz; Koenig, Hartmann;

Sauter, Hubert; Lorenz, Gisela; Ammermann, Eberhard

PATENT ASSIGNEE(S): BASF A.-G., Germany SOURCE: Eur. Pat. Appl., 86 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Fatent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

P.	ATENT NO.	KIND	DATE	APPLICATION NO.		DATE
– E	P 581095	A2	19940202	EP 1993-111103		19930712 <
	R: AT, BE, CH,	DE, DK	, ES, FR, GE	B, GR, IE, IT, LI,	NL, E	PT, SE
C.	A 2100546	A1	19940125	CA 1993-2100546		19930714 <
J	P 06211748	A	19940802	JP 1993-181305		19930722 <
A	U 9342121	A	19940127	AU 1993-42121		19930723 <
A	U 660226	B2	19950615			
Н	U 66105	A2	19940928	ни 1993-2150		19930723 <
Z.	A 9305332	A	19950123	ZA 1993-5332		19930723 <
PRIORI	TY APPLN. INFO.:			DE 1992-4224457	A	19920724
OTHER	SOURCE(S):	MARPAT	121:8893			
GT						

AB The title compds. [I; B = (un)substituted alkyl, C1-4 (un)substituted alkenyl, (un)substituted alkynyl, etc.; R1, R2 = (un)substituted alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkenyl, etc.; X, Y = H, halogen, CN, NO2, haloalkyl, alkyl, alkenyl, alkynyl, heteroaryl, heterocyclyl, etc.], useful as agrochem. fungicides, are prepared and I-containing formulations presented. Thus, Me α -(2-hydroxyphenyl)- β -methoxyacrylate was condensed with phenacyl bromide, producing acrylate II, m.p. 76°, which demonstrated 90% inhibitory activity against Plasmopara viticola at 250 ppm.

IT 154594-52-2P 154594-53-3P 154594-54-4P 154594-55-5P 154594-69-1P 154594-70-4P 154594-81-7P 154594-92-0P 154594-93-1P 154594-94-2P 154594-95-3P 154594-96-4P 154594-98-6P 154594-99-7P 154595-00-3P 154595-03-6P 154595-04-7P 154595-05-8P 154595-06-9P 154595-07-0P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as agrochem. fungicide)

RN 154594-52-2 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[(3-phenyl-5-isoxazolyl)methoxy]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 154594-53-3 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[[3-(2-methylphenyl)-5-isoxazolyl]methoxy]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 154594-54-4 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[[3-(3-methylphenyl)-5-isoxazolyl]methoxy]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

RN 154594-55-5 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[[3-(4-methylphenyl)-5-isoxazolyl]methoxy]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 154594-69-1 ZCAPLUS

CN Benzeneacetic acid, $2-[[5-(4-\text{chlorophenyl})-3-\text{isoxazolyl}]\text{methoxy}]-\alpha-$ (methoxymethylene)-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 154594-70-4 ZCAPLUS

CN Benzeneacetic acid, $2-[[3-(4-\text{chlorophenyl})-5-\text{isoxazolyl}]\text{methoxy}]-\alpha-$ (methoxymethylene)-, methyl ester, (E)- (9CI) (CA INDEX NAME)

RN 154594-81-7 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[1-(3-phenyl-5-isoxazolyl)ethoxy]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 154594-92-0 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[(5-phenyl-3-isoxazolyl)methoxy]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 154594-93-1 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[[5-(4-methylphenyl)-3-isoxazolyl]methoxy]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

RN 154594-94-2 ZCAPLUS

CN Benzeneacetic acid, 2-[(4-chloro-5-phenyl-3-isoxazolyl)methoxy]- α - (methoxymethylene)-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 154594-95-3 ZCAPLUS

CN Benzeneacetic acid, $2-[[5-(4-\text{chlorophenyl})-4-\text{methyl}-3-\text{isoxazolyl}]\text{methoxy}]-\alpha-(\text{methoxymethylene})-, \text{methyl ester, (E)- (9CI)}$ (CA INDEX NAME)

Double bond geometry as shown.

RN 154594-96-4 ZCAPLUS

CN Benzeneacetic acid, $2-[[5-(3-\text{chlorophenyl})-4-\text{methyl}-3-\text{isoxazolyl}]\text{methoxy}]-\alpha-(\text{methoxymethylene})-, \text{methyl ester, (E)- (9CI)}$ (CA INDEX NAME)

RN 154594-98-6 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[(1-phenyl-1H-pyrazol-4-yl)methoxy]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 154594-99-7 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[(5-methyl-1-phenyl-1H-pyrazol-4-yl)methoxy]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 154595-00-3 ZCAPLUS

CN Benzeneacetic acid, 2-[(3,5-dimethyl-1-phenyl-1H-pyrazol-4-yl)methoxy]- α -(methoxymethylene)-, methyl ester, (E)- (9CI) (CA INDEX NAME)

RN 154595-03-6 ZCAPLUS

CN Benzeneacetic acid, 2-[(4-chloro-3-cyclohexyl-5-isoxazolyl)methoxy]- α -(methoxymethylene)-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 154595-04-7 ZCAPLUS

CN Benzeneacetic acid, 2-[[4-chloro-3-(3-fluorophenyl)-5-isoxazolyl]methoxy]- $\alpha-(\text{methoxymethylene})-, \text{methyl ester, (E)- (9CI)}$ (CA INDEX NAME)

Double bond geometry as shown.

RN 154595-05-8 ZCAPLUS

CN Benzeneacetic acid, 2-[(4-ethyl-5-phenyl-3-isoxazolyl)methoxy]- α - (methoxymethylene)-, methyl ester, (E)- (9CI) (CA INDEX NAME)

154595-06-9 ZCAPLUS RN

CN Benzeneacetic acid, 2-[[4-chloro-5-(4-chlorophenyl)-3-isoxazolyl]methoxy]- α -(methoxymethylene)-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 154595-07-0 ZCAPLUS

CN Benzeneacetic acid, 2-[[4-chloro-5-(4-methylphenyl)-3-isoxazolyl]methoxy]- α -(methoxymethylene)-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

L89 ANSWER 69 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1994:284782 ZCAPLUS Full-text

DOCUMENT NUMBER: 120:284782

TITLE: Silver halide photographic material

INVENTOR(S): Takemura, Kumiko; Taguchi, Masaaki; Hashimoto,

Hiroyuki; Kawashima, Yasuhiko; Usaqawa, Yasushi; Inoe,

Kyoshi; Oohashi, Hirobumi

PATENT ASSIGNEE(S): Konishiroku Photo Ind, Japan Jpn. Kokai Tokkyo Koho, 72 pp. SOURCE:

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 05045790	A	19930226	JP 1991-201928	19910812 <
JP 3030578	B2	20000410		
PRIORITY APPLN. INFO.:			JP 1991-201928	19910812
GI				

In the title material comprising a support having thereon hydrophilic colloid layers (including one or more silver halide emulsion layers), at least one of said hydrophilic colloid layers contains a dispersion of solid microparticles of a dye compound represented by I, II, etc. For I, R1, R2 = substituent; R3, R4 = Ph ring having linking group connected to carboxyl group; L1 to L3 = methine; n = 0 to 2. For II, R1, R2 = substituent; R3, R4 = H, alkyl, cycloalkyl, alkenyl, etc.; L1 to L5 = methine; n, t = 0 or 1. At least one silver halide emulsion layer in the title material contains one or more 1-phenyl-5-mercaptotetrazole derivs. The title material shows high sensitivity and gives sharp images.

IT 150441-04-6

RL: TEM (Technical or engineered material use); USES (Uses) (photog. material containing)

RN 150441-04-6 ZCAPLUS

CN 1(2H)-Pyridineacetic acid, 3-[3-[3-(aminocarbonyl)-1-(2-carboxyphenyl)-5-hydroxy-1H-pyrazol-4-yl]-2-propenylidene]-5-cyano-3,6-dihydro-4-methyl-2,6-dioxo-(9CI) (CA INDEX NAME)

L89 ANSWER 70 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1994:245091 ZCAPLUS Full-text

DOCUMENT NUMBER: 120:245091

TITLE: Preparation of pyrazole containing propenoic ester

derivatives as agrochemical fungicides

INVENTOR(S): Hwang, Ki Jun; Kim, Sung Soo

PATENT ASSIGNEE(S): Korea Research Institute of Chemical Technology, S.

Korea

SOURCE: PCT Int. Appl., 37 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

GΙ

PA.	PATENT NO.				KIND DATE			APPLICATION NO.					DATE					
WO	9400	436			A1	_	 1994	 0106		 WO 1	 993-:	KR52			1	9930	623	<
	W:	ΑT,	ΑU,	BB,	BG,	BR,	CA,	CH,	CZ,	DE,	DK,	ES,	FI,	GB,	HU,	JP,	KP,	
		LK,	LU,	MG,	MN,	MW,	NL,	NO,	NΖ,	PL,	PT,	RO,	RU,	SD,	SE,	SK,	UA,	US
	RW:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,	
		BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	ML,	MR,	NE,	SN,	TD,	ΤG			
KR	9506	150			В1		1995	0609		KR 1	992-	1115	0		1	9920	625	<
AU	9454	187			Α		1994	0124		AU 1	994-	5418	7		1	9930	623	<
PRIORIT	Y APP	LN.	INFO	.:						KR 1	992-	1115	0		A 1	9920	625	
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OTHER SO	OURCE	(S):			MAR:	PAT	120:	2450	91									

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CH₂O

R₁

R₂

- AB Title compds. I (R = H, one or more halo, Me, alkyl, alkoxy, O2N, Ph; R1 = Me, alkyl, alkenyl, alkynyl, PhCH2, aryl, (substituted), pyridyl; R2, R3 = H, halo, F3C, haloalkyl; X = C, N) are prepared. To Ph3P+CH2OMe Br- in THF was added EtCHMeLi in cyclohexane followed by Me 2-[2-[[1-methyl-5-(trifluoromethyl)-3-pyrazolyl]methyl]phenyl]glyoxylate in THF to give I (R = R2 = H, R1 = Me. R3 = F3C, X = trans-CH) which showed EC50 against wheat leaf rust and barley powdery mildew of <0.4 and <0.08 ppm, resp.
- IT 154315-23-8P 154315-24-9P 154315-25-0F

 RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as agrochem. fungicide)
- RN 154315-23-8 ZCAPLUS
- CN Benzeneacetic acid, α -(methoxymethylene)-2-[[[1-phenyl-3-(trifluoromethyl)-1H-pyrazol-5-yl]oxy]methyl]-, methyl ester, (Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

- RN 154315-24-9 ZCAPLUS
- CN Benzeneacetic acid, α -(methoxymethylene)-2-[[[1-phenyl-3-(trifluoromethyl)-1H-pyrazol-5-yl]oxy]methyl]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

- RN 154315-25-0 ZCAPLUS
- CN Benzeneacetic acid, α -(methoxyimino)-2-[[[1-(2-pyridinyl)-3-(trifluoromethyl)-1H-pyrazol-5-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

IT 154315-40-9 154315-41-0

RL: RCT (Reactant); RACT (Reactant or reagent)

(reaction of, in preparation of agrochem. fungicides)

RN 154315-40-9 ZCAPLUS

CN Benzeneacetic acid, α -oxo-2-[[[1-phenyl-3-(trifluoromethyl)-1H-pyrazol-5-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 154315-41-0 ZCAPLUS

CN Benzeneacetic acid, α -oxo-2-[[[1-(2-pyridinyl)-3-(trifluoromethyl)-1H-pyrazol-5-yl]oxy]methyl]-, methyl ester (9CI) (CA INDEX NAME)

L89 ANSWER 71 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1994:244258 ZCAPLUS Full-text

DOCUMENT NUMBER: 120:244258

TITLE: A short synthesis of potential juvenoids based on the

isoxazole chemistry

AUTHOR(S): Martin, Lourdes; Polo, Cecilia; Ramos, Vicente;

Torroba, Tomas; Marcaccini, Stefano

CORPORATE SOURCE: Fac. Vet., Univ. Extremadura, Caceres, 10071, Spain

SOURCE: Heterocycles (1993), 36(10), 2259-65

CODEN: HTCYAM; ISSN: 0385-5414

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 120:244258

GΙ

3,4,5-Trisubstituted isoxazoles 1 (shown as I) and 4 (shown as II) afforded, after chromic oxidation and borohydride reduction, (\pm) -3-methyl-6-(3- methyl-5-phenylisoxazol-4-yl)-6-hydroxyhexanoic acid (2) or (\pm) -1-(3-methyl-5-phenylisoxazol-4-yl)-3,4-dihydro-1H-2-benzopyran-3- one (5) which were reduced to (\pm) - (\mathbb{Z}/\mathbb{E}) -3-methyl-7-benzoyl-8-oxonon-6- enoic acid (3) and (\mathbb{E}) -2-(2-(2-benzoyl-3-oxobut-1-enyl]phenyl)acetic acid (6) with molybdenum hexacarbonyl. Lactone (5) afforded a single E-diastereoisomer of acid (6). Catalytic hydrogenation of 5 afforded selectively an isoxazole which was reduced with molybdenum hexacarbonyl to 2-(2-[2-benzoyl-3-oxobutyl]phenyl)acetic acid (8). Structures of products are related with those of some juvenoids.

IT 154051-10-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reduction of, with molybdenum hexacarbonyl)

RN 154051-10-2 ZCAPLUS

CN Benzeneacetic acid, 2-[(3-methyl-5-phenyl-4-isoxazolyl)methyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{N} \\ \text{Ph} \\ \text{CH}_2 \\ \text{CH}_2 - \text{CO}_2 \text{H} \end{array}$$

L89 ANSWER 72 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1994:106997 ZCAPLUS Full-text

DOCUMENT NUMBER: 120:106997

TITLE: Preparation of pyrazole derivatives and agrochemical

fungicides

INVENTOR(S): Kasahara, Isamu; Iihama, Teruyuki; Sugiura, Tadashi;

Hashimoto, Sho; Sano, Shinsuke; Hosokawa, Hiroyasu;

Yokota, Chinami

PATENT ASSIGNEE(S): Nippon Soda Co., Ltd., Japan

SOURCE: PCT Int. Appl., 90 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Fatent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.					KIND DATE			,	APPLICATION NO.				DATE					
	WO	9307	 138			A1	_	 1993	 0415		 WO 1	 992-	JP13	03		19	 9921	007	<
		W:	ΑT,	ΑU,	BB,	BG,	BR,	CA,	CH,	CS,	DE,	DK,	ES,	FΙ,	GB,	HU,	JP,	KR,	
			LK,	LU,	MG,	MN,	MW,	NL,	NO,	PL,	RO,	RU,	SD,	SE,	US				
		RW:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	SE,	BF,	
			ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	ML,	MR,	SN,	TD,	TG					
	AU	9226	970			Α		1993	0503		AU 1	992-	2697	0		19	9921	007	<
	CN	1071	424			Α		1993	0428		CN 1	992-	1112	27		19	9921	800	<
PRI	ORIT	APP:	LN.	INFO	.:						JP 1	991-	2891.	58		A 19	9911	800	
											JP 1	992-	1315	71		A 19	9920	424	
											JP 1	992-	1974	57		A 19	9920	702	
											WO 1	992-	JP13	03		A 19	9921	007	

OTHER SOURCE(S): MARPAT 120:106997

$$A-B-Q$$
 Y
 R^{1}
 R^{2}

GΙ

The title compds. [I; Y = CR6, N; R1, R2, R3, R4, R6 = H, halo, (un)substituted alkyl, (un)substituted alkoxy, (un)substituted alkenyloxy, etc.; R5 = H, halo, (un)substituted alkyl, (un)substituted alkoxy, etc.; A = (un)substituted aryl, (un)substituted heterocyclyl; B = (un)substituted alkylene, etc.; Q = (un)substituted pyrazolediyl] are prepared E.g., Et 4-(4-chlorophenyl)-3-oxobutanoate in EtOH was refluxed with (6-methyl-2-pyridyl)hydrazine to give 1-(6-methyl-2-pyridyl)-3-(4-chlorobenzyl)-5-hydroxy-1H-pyrazole, which was O-methylated with MeI to give 1-(6-methyl-2-pyridyl)-3-(4-chlorobenzyl)-5-methoxy-1H-pyrazole. This at 200 ppm effected >90% kill of Cercospora beticola.

IT 150400-56-9P 150400-57-0P 150400-58-1P

150400-61-6P 150400-78-5P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as agrochem. fungicide)

RN 150400-56-9 ZCAPLUS

CN Benzoic acid, 4-[[5-methoxy-1-(6-methyl-2-pyridinyl)-1H-pyrazol-3-yl]methyl]- (9CI) (CA INDEX NAME)

RN 150400-57-0 ZCAPLUS

CN Benzoic acid, 4-[[5-methoxy-1-(6-methyl-2-pyridinyl)-1H-pyrazol-3-yl]methyl]-, methyl ester (9CI) (CA INDEX NAME)

$$MeO-C$$

$$OMe$$

$$N$$

$$OMe$$

RN 150400-58-1 ZCAPLUS

CN Benzamide, 4-[[5-methoxy-1-(6-methyl-2-pyridinyl)-1H-pyrazol-3-yl]methyl]-(9CI) (CA INDEX NAME)

$$H_2N-C$$

$$OMe$$

$$OMe$$

RN 150400-61-6 ZCAPLUS

CN Carbamic acid, [4-[[5-methoxy-1-(6-methyl-2-pyridinyl)-1H-pyrazol-3-yl]methyl]phenyl]-, ethyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{CH}_2 \\ \text{NMe} \end{array}$$

RN 150400-78-5 ZCAPLUS

CN Benzoic acid, 4-[[5-methoxy-4-methyl-1-(6-methyl-2-pyridinyl)-1H-pyrazol-3-yl]methyl]-, methyl ester (9CI) (CA INDEX NAME)

$$MeO-C$$

$$MeO-$$

ACCESSION NUMBER: 1994:106561 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 120:106561

TITLE: Preparation of carbamates and plant-protecting agents

containing them

INVENTOR(S): Mueller, Bernd; Sauter, Hubert; Roehl, Franz; Doetzer,

Reinhard; Lorenz, Gisela; Ammermann, Eberhard

PATENT ASSIGNEE(S): BASF A.-G., Germany

SOURCE: PCT Int. Appl., 764 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Fatent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

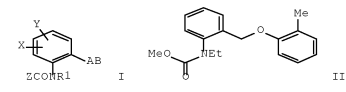
PATENT INFORMATION:

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	9315 W:	046 AT, LU,	AU, MG,	BG, MN,	A1 BR, MW,	CA,	1993 , CH, , NO,	0805 DE, PL,	DK, RO,	VO 1 ES, RU	993- FI,	EP1(GB,)4 HU,	JP,	19 KP,	9930: KR,	118 LK,	
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	4234 4234				A1		1994 1994 1994 1994	0414	L	DE I	992-	4234	1020		13	99ZI	109	<
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	4234				A1		1994	0414	L	ノ比 I フロ 1	992-	4234	1001		10	99ZI	109	<
					AΙ		1994	0001	L 7	ノĽ 1	992-	2251	1081 L4		1:	39ZI	110	<
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	2116	136 10			T3		1998	0716					227					
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	1044				A		2003	0 / 2 1		лд <u>т</u> гт 1	332-	.104/	7110 189		1 (3330. 3030:	122	/
	9300				A		1994		7	LЫ Т 7Л 1	332-	.604	109		10	3330. 3330.	122	\
	9403				A		1994						3					
	9402				A		1994						1			9940		
	3024						1998			10 I	J J 4	2015	İ		Δ.) J 4 U	120	
	5824	705			B1 A		1998			IC 1	991_	.2566	528		1 (aa an	729	/
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US 1994-256628 A1 19940729 US 1998-110884 A3 19980707 US 1999-275767 A3 19990325

OTHER SOURCE(S):

CASREACT 120:106561; MARPAT 120:106561



Title compds. [I; Z = MeO, NH2, NHMe, NMe2, Me, Et, CF3, CCl3; X, Y = H, F, AΒ Cl, Br, cyano, NO2, alkoxy, alkenyloxy, alkynyloxy, alkyl, alkenyl, alkynyl; XY = atoms to form a (substituted) (hetero)aromatic, alicyclic, heterocyclic, partially or fully hydrogenated ring; R1 = H, (substituted) alkyl, alkenyl, alkynyl, cyclopropyl, cyclopropylmethyl, cyclobutyl, CH2CN, CH2OMe, CO2Me, alkoxy, alkenyloxy, alkynyloxy, etc.; A = 0, S, CR2:NO, C.tplbond.C, CHR2O2C, OCHR2, bond, etc.; R2 = H, alkyl, alkenyl, alkynyl, cycloalkyl; B = H, cycloalkyl, cycloalkenyl, cycloalkynyl, aryl, heteroaryl, heterocyclyl, arylalkyl, etc.], were prepared Thus, o-toluidine was stirred with ClCO2Me in CH2Cl2 to give 100% 2-MeC6H4NHCO2Me, which in DMF was treated with NaH and EtI to give 93% 2-MeC6H4NEtCO2Me. This was irradiated with NBS and azobisisobutyronitrile in CC14 using a 300 W UV lamp to give 2-BrCH2C6H4NEtCO2Me. This was stirred with p-cresol and NaH in DMF to give title compound II. Numerous I as 25 ppm sprays gave 95% control of Erysiphe graminis on wheat.

IT 151828-02-3P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as agrochem. fungicide)

RN 151828-02-3 ZCAPLUS

CN Carbamic acid, methoxy[2-[[(1-phenyl-1H-pyrazol-4-yl)oxy]methyl]phenyl]-, methyl ester (9CI) (CA INDEX NAME)

ACCESSION NUMBER: 1993:539098 ZCAPLUS Full-text

DOCUMENT NUMBER: 119:139098

TITLE: Preparation of dihydropyran derivatives and plant

protecting agents containing them

INVENTOR(S): Mueller, Bernd; Brand, Siegbert; Sauter, Hubert;

Roehl, Franz; Ammermann, Eberhard; Lorenz, Gisela

PATENT ASSIGNEE(S): BASF A.-G., Germany

SOURCE: Eur. Pat. Appl., 153 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Fatent
LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

GΙ

PATENT NO.		KIND	DATE	APPLICATION NO.		DATE	
EP 534216		A1	19930331	EP 1992-115247		19920905	<
EP 534216		B1	19980819				
R: AI	, BE, CH,	DE, DK	, ES, FR,	GB, GR, IT, LI, NL, E	PT, SE		
DE 4131311		A1	19930401	DE 1991-4131311		19910920	<
JP 0521392	8	A	19930824	JP 1992-232502		19920831	<
AT 169911		Τ	19980915	AT 1992-115247		19920905	<
IL 103157		A	19980222	IL 1992-103157		19920914	<
CA 2078625		A1	19930321	CA 1992-2078625		19920918	<
AU 9224566		A	19930325	AU 1992-24566		19920918	<
AU 651003		B2	19940707				
HU 61879		A2	19930329	ни 1992-2996		19920918	<
HU 213029		В	19970128				
ZA 9207152		A	19940318	ZA 1992-7152		19920918	<
US 5536734		A	19960716	US 1994-263414		19940621	<
PRIORITY APPLN.	INFO.:			DE 1991-4131311	A	19910920	
				US 1992-946651	В1	19920918	
OTHER SOURCE(S)	:	MARPAT	119:13909	98			

$$R^1$$
 R^2
 R^3
 R^4
 R^4
 R^3
 R^4
 R^4
 R^3
 R^4
 Title compds. [I; U = CHOR5, CHSR5, CH2, CHR5, CHX, NOR5; X = halo; A = bond, CHR6, (CHR7CHR6)n, (CR21:CHR20)mCR7:CR6, C.tplbond.C, OCHR6, SCHR6, NR18CHR6, CO2CHR6, R19C:NOCHR6; B = H, (substituted) alkyl, alkenyl, alkynyl, cycloalkyl, (hetero)aryl, heterocyclyl, cycloalkenyl; R1 = H, OR8, (substituted) aryloxy; R2 = R9, (substituted) aryl; R3 = R10, (substituted) aryl, CHR11OR12, CO2R12, CONR12R13, CHR11CHR14B; R4 = OR15, NR16R17, R25; n = 1-3; m = 0, 1; R5, R8, R12, R13, R18, R25 = (substituted) alkyl, alkenyl, alkynyl, cycloalkyl; R6, R7, R11, R16, R17, R20, R21 = H, R5; R19 = H, cyano, (substituted) (cyclo)alkyl; R9, R10 = H, (substituted) (cyclo)alkyl, alkynyl;

with provisos], were prepared Thus, 6-cyano-2,3-dihydropyran was reduced with DIBAL to give 78% 6-formyl-2,3-dihydropyran, which was treated with (PhCH2)Ph3PCl/KOCMe3 in THF to give 85% 6-phenethenyl-2,3-dihydropyran. This was treated with Me oxalate and pyridine in CH2Cl2 to give 94% Me 6-trans-phenethenyl-2,3- dihydropyranyl-5-glyoxalate. This was treated with EtPh3PCl/KOCMe3 in THF to give 37% title compound II. Numerous I exhibited 95% control of Plasmopara viticola on grapevines. I are also said to be insecticides, nematocides, and plant growth regulators.

IT 149795-21-1F 149795-22-2F 149795-95-9F
RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as agrochem.)

RN 149795-21-1 ZCAPLUS

CN 2H-Pyran-5-acetic acid, 6-[2-[3-(4-chlorophenyl)-5-isoxazolyl] ethenyl]-3,4-dihydro- α -(methoxymethylene)-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

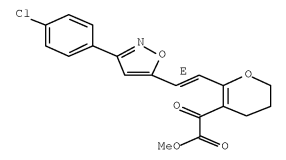
RN 149795-22-2 ZCAPLUS

CN 2H-Pyran-5-acetic acid, 6-[2-[3-(4-chlorophenyl)-5-isoxazolyl] ethenyl]-3,4-dihydro- α -(methoxymethylene)-, methyl ester, (Z,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 149795-95-9 ZCAPLUS

CN 2H-Pyran-5-acetic acid, $6-[2-[3-(4-\text{chlorophenyl})-5-\text{isoxazolyl}] = \text{thenyl}]-3, 4-dihydro-α-oxo-, methyl ester, (E)- (9CI) (CA INDEX NAME)$



L89 ANSWER 75 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1993:528317 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 119:128317

TITLE: Silver halide photographic material with good

decolorization

INVENTOR(S): Yamada, Taketoshi; Hanyu, Takeshi PATENT ASSIGNEE(S): Konishiroku Photo Ind, Japan SOURCE: Jpn. Kokai Tokkyo Koho, 22 pp.

CODEN: JKXXAF

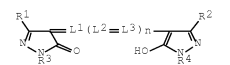
DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

GI

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 05045787 PRIORITY APPLN. INFO.:	А	19930226	JP 1991-200510 JP 1991-200510	19910809 < 19910809
OTHER SOURCE(S):	MARPAT	119:128317	01 1331 200010	19910009



- AB The title material has a photog. constituent layer containing a dispersion of particles of a dye represented, e.g., by I. For I, R1, R2 = CO2H or substituent having CO2H; R3, R4 = H or substituent which has no CO2H; L1-L3 = methine; n = 0 to 2. The above-mentioned photog. constituent layer is located on a photosensitive silver halide emulsion layer which contains an organic compound which reacts with the developing agent. The title material shows good decolorization after photog. processing.
- IT 149489-71-4

RL: TEM (Technical or engineered material use); USES (Uses) (photog. materials containing)

RN 149489-71-4 ZCAPLUS

CN 1,4-Benzenedicarboxylic acid, 2-[4-[3-[1-(carboxymethyl)-5-cyano-1,6-dihydro-4-methyl-2,6-dioxo-3(2H)-pyridinylidene]-1-propenyl]-3-cyano-5-

$$CO_2H$$
 NC
 CH
 L89 ANSWER 76 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1993:428146 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 119:28146

TITLE: Preparation of α -(azolylvinylaryl)- β -

methoxyacrylates as pesticides

INVENTOR(S): Kirstgen, Reinhard; Theobald, Hans; Koenig, Hartmann;

Harreus, Albrecht; Oberdorf, Klaus; Kardorff, Uwe; Harries, Volker; Lorenz, Gisela; Ammermann, Eberhard

PATENT ASSIGNEE(S): BASF A.-G., Germany SOURCE: Ger. Offen., 29 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Fatent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NC	. K	IND	DATE	APPLICATION NO.		DATE	
DE 412699	4	 A1	19930218	DE 1991-4126994		19910816	<
JP 052138	67	A	19930824	JP 1992-204996		19920731	<
JP 321490	6	В2	20011002				
EP 528245		A1	19930224	EP 1992-113197		19920803	<
EP 528245		В1	19971112				
R: A	T, BE, CH, D	E, DK,	ES, FR,	GB, GR, IT, LI, NL,	PT,	SE	
AT 160142		T	19971115	AT 1992-113197		19920803	<
ES 211045	6	Т3	19980216	ES 1992-113197		19920803	<
IL 102729		A	19981030	IL 1992-102729		19920804	<
CA 207541	6	A1	19930217	CA 1992-2075416		19920806	<
US 540383	8 .	A	19950404	US 1992-928038		19920811	<
AU 922100	5 .	A	19930218	AU 1992-21005		19920814	<
AU 648193		В2	19940414				
HU 61652		A2	19930301	HU 1992-2653		19920814	<
HU 212604		В	19960930				
ZA 920612	0 .	A	19940214	ZA 1992-6120		19920814	<
KR 221506		В1	19990915	KR 1992-14661		19920814	<
PRIORITY APPLN	. INFO.:			DE 1991-4126994	А	19910816	

OTHER SOURCE(S):

CASREACT 119:28146; MARPAT 119:28146

Title compds. [I; X = C, N; Y, Z = CR4, N, O, S; n = 0-4; R1 = NO2, cyano, halo, (halo)alkyl, (halo)alkoxy, alkylthio; (R1)2 = (substituted) 1,3-butadien-1,4-diyl; R2 = (halo)alkyl, halo, cyano, NO2, alkoxycarbonyl, Me2N, H; R3 = H, (substituted) alkyl, (substituted) (saturated) (O-, S-, or N-containing) ring system, (substituted) mono- or bicyclic aryl; R4 = H, (halo)alkyl, halo, cyano, NO2, Me2N, alkoxycarbonyl], were prepared as pesticides (no data). Thus, di-Me 2-(β -methoxy-2-methoxycarbonylvinyl)benzylphosphonate and 5-methyl-1-phenylpyrazol-4-ylcarboxaldehyde were stirred with NaH in THF overnight to give title compound II.

CN Benzeneacetic acid, 2-[2-[1-(4-chlorophenyl)-3-methyl-1H-pyrazol-4-yl]ethenyl]- α -(methoxymethylene)-, methyl ester, (E,?)- (9CI) (CA INDEX NAME)

Double bond geometry as described by E or Z.

RN 148001-22-3 ZCAPLUS

CN Benzeneacetic acid, $2-[2-[1-(4-\text{chloropheny1})-3,5-\text{dimethyl-1H-pyrazol-4-yl]ethenyl}]-\alpha-(methoxymethylene)-, methyl ester, (E,?)- (9CI) (CA INDEX NAME)$

Double bond geometry as described by ${\tt E}$ or ${\tt Z}$.

RN 148001-23-4 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[2-(5-methyl-1-phenyl-1H-pyrazol-4-yl)ethenyl]-, methyl ester, (E,?)- (9CI) (CA INDEX NAME)

Double bond geometry as described by ${\tt E}$ or ${\tt Z}$.

RN 148001-24-5 ZCAPLUS

CN Benzeneacetic acid, 2-[2-[1-(4-chlorophenyl)-5-methyl-1H-pyrazol-4-yl]ethenyl]- α -(methoxymethylene)-, methyl ester, (E,?)- (9CI) (CA INDEX NAME)

Double bond geometry as described by E or Z.

RN 148001-26-7 ZCAPLUS

CN Benzeneacetic acid, 2-[2-[4-chloro-3-phenyl-5-isoxazolyl]ethenyl]- α - (methoxymethylene)-, methyl ester, (E,?)- (9CI) (CA INDEX NAME)

Double bond geometry as described by E or Z.

RN 148001-27-8 ZCAPLUS

CN Benzeneacetic acid, 2-[2-[4-chloro-3-(4-methylphenyl)-5-isoxazolyl]ethenyl]- α -(methoxymethylene)-, methyl ester, (E,?)-(9CI) (CA INDEX NAME)

Double bond geometry as described by E or Z.

RN 148001-28-9 ZCAPLUS

CN Benzeneacetic acid, 2-[2-[4-chloro-3-(4-fluorophenyl)-5-isoxazolyl]ethenyl]- α -(methoxymethylene)-, methyl ester, (E,?)-(9CI) (CA INDEX NAME)

Double bond geometry as described by E or Z.

RN 148001-29-0 ZCAPLUS

CN Benzeneacetic acid, 2-[2-[4-chloro-3-(3,4-difluorophenyl)-5-isoxazolyl]ethenyl]- α -(methoxymethylene)-, methyl ester, (E,?)-(9CI) (CA INDEX NAME)

Double bond geometry as described by E or Z.

RN 148001-30-3 ZCAPLUS

CN Benzeneacetic acid, 2-[2-[4-chloro-3-(2,6-difluorophenyl)-5-isoxazolyl]ethenyl $]-\alpha-(\text{methoxymethylene})-,$ methyl ester, (E,?)-(9CI) (CA INDEX NAME)

Double bond geometry as described by E or Z.

RN 148001-31-4 ZCAPLUS

CN Benzeneacetic acid, 2-[2-[4-chloro-3-(4-chlorophenyl)-5-isoxazolyl]ethenyl]- α -(methoxymethylene)-, methyl ester, (E,?)-

(9CI) (CA INDEX NAME)

Double bond geometry as described by E or Z.

RN 148001-32-5 ZCAPLUS

CN Benzeneacetic acid, 2-[2-[4-chloro-3-(3-chloropheny1)-5-isoxazolyl]ethenyl]- α -(methoxymethylene)-, methyl ester, (E,?)-(9CI) (CA INDEX NAME)

Double bond geometry as described by E or Z.

RN 148001-33-6 ZCAPLUS

CN Benzeneacetic acid, $2-[2-(4-\text{ethyl}-5-\text{phenyl}-3-\text{isoxazolyl})\text{ethenyl}]-\alpha-$ (methoxymethylene)-, methyl ester, (E,?)- (9CI) (CA INDEX NAME)

Double bond geometry as described by E or Z.

RN 148001-34-7 ZCAPLUS

CN Benzeneacetic acid, $2-[2-[5-(4-\text{chlorophenyl})-4-\text{ethyl}-3-\text{isoxazolyl}]\text{ethenyl}]-\alpha-(\text{methoxymethylene})-, methyl ester, (E,?)- (9CI) (CA INDEX NAME)$

Double bond geometry as described by E or Z.

RN 148001-35-8 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[2-[3'-(1-methylethyl)[3,5'-biisoxazol]-5-yl]ethenyl]-, methyl ester, (E,?)- (9CI) (CA INDEX NAME)

Double bond geometry as described by E or Z.

RN 148001-36-9 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[2-[3-(1-methyl-1H-pyrazol-3-yl)-5-isoxazolyl]ethenyl]-, methyl ester, (E,?)- (9CI) (CA INDEX NAME)

Double bond geometry as described by E or Z.

L89 ANSWER 77 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1993:428133 ZCAPLUS Full-text

DOCUMENT NUMBER: 119:28133

TITLE: Derivatives of β -substituted cinnamic acid

INVENTOR(S): Sauter, Hubert; Oberdorf, Klaus; Wingert, Horst; Von

Deyn, Wolfgang; Grammenos, Wassilios; Koenig, Hartmann; Rang, Harald; Roehl, Franz; et al.

PATENT ASSIGNEE(S): BASF A.-G., Germany

SOURCE: Eur. Pat. Appl., 127 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PAT	TENT NO.			KINI)	DATE		А	PF	PLICATION NO.			DATE	
		525516			A2	_		0203	– E	P	1992-112086			19920715	<
		525516			A3		1993								
	EP	525516			В1		1995								
			BE,	CH,		DK,					R, IT, LI, NL,				
	DΕ	4124989			A1		1993	0204	D	E	1991-4124989			19910727	<
	ΑT	128454			Τ		1995	1015	А	Τ.	1992-112086			19920715	<
	ES	2078602			Т3		1995	1216	E	S	1992-112086			19920715	<
	JΡ	05255191	_		А		1993	1005	J	Ρ	1992-190680			19920717	<
	HU	61519			A2		1993	0128	Н	U	1992-2451			19920724	<
	HU	213456			В		1997	0630							
	AU	9220590			Α		1993	0128	А	U	1992-20590			19920727	<
	AU	653612			В2		1994	1006							
	ZA	9205613			Α		1994	0127	Z	Α	1992-5613			19920727	<
	CA	2075354			A1		1993	0128	С	Α	1992-2075354			19920803	<
	US	5538940			А		1996	0723	U	S	1995-440126			19950512	<
	US	5573999			А		1996	1112	U	S	1995-441639			19950515	<
PRIOF	RITY	APPLN.	INFO	. :					D	E	1991-4124989	I	Ā	19910727	
									[]	I.S	1992-919270	F	31	19920727	
											1993-173936			19931228	
									0	\sim	1000 110000		-	10001220	

GΙ

AB Title compds. (235 compds.) were prepared as inhibitors of mitochondrial respiration. Thus, 2-MeC6H4Ac was treated with (MeO)2CO to give 94% 2-MeC6H4COCH2CO2Me which was enol methylated to give 94% (E)-2-MeC6H4C(OMe):CHCO2Me. The latter compound was brominated, oxidized to the aldehyde, and treated with 2-(4-fluorophenyl)-4- thiazolylmethylphosphonium chloride to give the cinnamate I. At 1.8 + 10-5 mol/L I caused 96 and 99% inhibition of mitochondrial respiration in Saccharomyces cerevisiae and Musca domestica resp.

IT 147500-08-1P 147500-09-2P 147500-10-5P 147500-11-6P 147500-12-7P 147500-13-8P

147500-14-9P 147500-15-0P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and fungicidal activity of)

RN 147500-08-1 ZCAPLUS

CN 2-Propenoic acid, 3-[2-[2-[1-(3-chlorophenyl)-1H-pyrazol-4-

yl]ethenyl]phenyl]-3-methoxy-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 147500-09-2 ZCAPLUS

CN 2-Propenoic acid, 3-[2-[2-[1-(3-chlorophenyl)-1H-pyrazol-4-yl]ethenyl]-3-methoxy-, methyl ester, (E,Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 147500-10-5 ZCAPLUS

CN 2-Propenoic acid, 3-methoxy-3-[2-[2-[1-(4-methoxyphenyl)-1H-pyrazol-4-yl]ethenyl]phenyl]-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 147500-11-6 ZCAPLUS

CN 2-Propenoic acid, 3-methoxy-3-[2-[2-[1-(4-methoxyphenyl)-1H-pyrazol-4-

yl]ethenyl]phenyl]-, methyl ester, (E,Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 147500-12-7 ZCAPLUS

CN 2-Propenoic acid, 3-[2-[2-[1-(4-chlorophenyl)-1H-pyrazol-4-yl]ethenyl]phenyl]-3-methoxy-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 147500-13-8 ZCAPLUS

CN 2-Propenoic acid, 3-[2-[2-[1-(4-chlorophenyl)-1H-pyrazol-4-yl]ethenyl]-3-methoxy-, methyl ester, (E,Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 147500-14-9 ZCAPLUS

CN 2-Propenoic acid, 3-[2-[2-[5-(4-chlorophenyl)-4-methyl-3-isoxazolyl]ethenyl]phenyl]-3-methoxy-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 147500-15-0 ZCAPLUS

CN 2-Propenoic acid, 3-[2-[2-[5-(4-chlorophenyl)-4-methyl-3-isoxazolyl]ethenyl]phenyl]-3-methoxy-, methyl ester, (E,Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

IT 147499-97-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and fungicidal and insecticidal activity of)

RN 147499-97-6 ZCAPLUS

CN 2-Propenoic acid, 3-[2-[3-(4-chlorophenyl)-5-isoxazolyl]ethenyl]-3-methoxy-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

IT 147500-53-6P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 147500-53-6 ZCAPLUS

CN 2-Propenoic acid, 3-methoxy-3-[2-[2-[3-(6-methyl-2-pyridinyl)-5-isoxazolyl]ethenyl]phenyl]-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

L89 ANSWER 78 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1993:124559 ZCAPLUS Full-text

DOCUMENT NUMBER: 118:124559

TITLE: Preparation of (heterocyclyl)- α -phenylacrylates

as agrochemical fungicides

INVENTOR(S): Grammenos, Wassilios; Kirstgen, Reinhard; Oberdorf,

Klaus; Sauter, Hubert; Roehl, Franz; Otter, Rainer; Ammermann, Eberhard; Lorenz, Gisela; Kardorff, Uwe;

Kuenast, Christoph

PATENT ASSIGNEE(S): BASF A.-G., Germany

SOURCE: Eur. Pat. Appl., 190 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

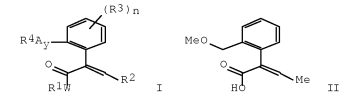
PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE	
EP 513580 A2 19921119 EP 1992-107059 19920424	<
EP 513580 A3 19930331	
EP 513580 B1 19961023	
R: AT, BE, CH, DE, DK, ES, FR, GB, IT, LI, NL, PT, SE	
DE 4116090 A1 19921119 DE 1991-4116090 19910517	<
AT 144502 T 19961115 AT 1992-107059 19920424	<
ES 2094842 T3 19970201 ES 1992-107059 19920424	<
JP 05213815 A 19930824 JP 1992-111088 19920430	<

JP 3234274	В2	20011204		
IL 101740	A	19970610	IL 1992-101740	19920430 <
CA 2068017	A1	19921118	CA 1992-2068017	19920505 <
AU 9216268	A	19921119	AU 1992-16268	19920515 <
AU 648664	B2	19940428		
HU 61435	A2	19930128	HU 1992-1631	19920515 <
HU 213444	В	19970630		
ZA 9203534	А	19931115	ZA 1992-3534	19920515 <
KR 201241	B1	19990615	KR 1992-8243	19920515 <
US 5298527	A	19940329	US 1993-103154	19930809 <
US 5416068	A	19950516	US 1994-176649	19940103 <
PRIORITY APPLN. INFO.:			DE 1991-4116090	A 19910517
			US 1992-878295 E	B1 19920506
			US 1993-103154 A	A3 19930809

OTHER SOURCE(S): MARPAT 118:124559

GΙ



AB Title compds. [I; n = 0-4; yl = 0, 1; R1 = H, (halo-substituted) alkyl, alkenyl, alkynyl, cycloalkyl, vinyl, ethynyl; R2 = cyano, alkenyl, alkynyl, (substituted) cycloalkyl, heterocyclyl, alkyl; R3 = H, NO2, cyano, halo, (halo)alkyl, (halo)alkoxy, (halo)alkylthio; 2 adjacent R3's = R4 = H, CHO, (substituted) alkyl, alkenyl, alkynyl, (unsatd.) carbocyclyl, heterocyclyl, aryl, etc.; W = bond, O, S, imino; A = O, CO, O2C, S, SO, SO2, alkenylene, alkynylene, alkylene, imino, carbonylimino, N:N, etc.], were prepared Thus, Ph3PEtBr, Me 2-methoxymethylphenylglyoxylate (preparation given), and KOCMe3 were stirred in THF at 5-25° to give a mixture of olefins which was saponified with aqueous KOH to give title compound II. Numerous I as 250 ppm sprays reduced infestation of grape plants by Plasmopara viticola to 0-15%, vs. 70% for untreated controls.

IT 145911-86-0P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of as agrochem. fungicide)

RN 145911-86-0 ZCAPLUS

CN Benzeneacetic acid, $2-[2-[3-(3-\text{chloropheny1})-5-\text{isoxazoly1}] = \alpha - \text{ethylidene-, methyl ester, (E,Z)- (9CI) (CA INDEX NAME)}$

Double bond geometry as shown.

```
RN 145910-27-6 ZCAPLUS CN Benzeneacetic acid, \alpha-ethylidene-2-[[(5-phenyl-3-isoxazolyl)oxy]methyl]-, methyl ester, (E)- (9CI) (CA INDEX NAME)
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RN 145910-52-7 ZCAPLUS CN Benzeneacetic acid, 2-[2-chloro-2-[3-(3-chlorophenyl)-5-isoxazolyl]ethenyl]- α -ethylidene-, methyl ester, (Z,E)- (9CI) (CFINDEX NAME)

Double bond geometry as shown.

$$\begin{array}{c|c} & & & & \\ \hline \\ C1 & & & \\ \hline \\ C1 & & \\ \hline \\ C1 & & \\ \hline \\ E & \\ Me \\ \hline \end{array}$$

RN 145910-53-8 ZCAPLUS CN Benzeneacetic acid, 2-[2-chloro-2-[3-(3-chlorophenyl)-5-isoxazolyl]ethenyl]- α -ethylidene-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 145910-54-9 ZCAPLUS

CN Benzeneacetic acid, 2-[2-[3-(3-chlorophenyl)-5-isoxazolyl]ethenyl]- α -ethylidene-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

RN 145910-55-0 ZCAPLUS CN Benzeneacetic acid, 2-[2-chloro-2-[3-(3-chlorophenyl)-5-isoxazolyl]ethenyl]- α -propylidene-, methyl ester, (Z,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & &$$

RN 145910-56-1 ZCAPLUS CN Benzeneacetic acid, 2-[2-chloro-2-[3-(3-chlorophenyl)-5-isoxazolyl]ethenyl]- α -propylidene-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

$$\begin{array}{c|c} & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & \\ & & \\$$

RN 145910-64-1 ZCAPLUS CN Benzeneacetic acid, 2-[[[1-(2,4-dichlorophenyl)-1H-pyrazol-4-yl]oxy]methyl]- α -ethylidene-, methyl ester, (E)- (9CI) (CA INDEX NAME)

RN 145911-06-4 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[2-[3-(3-fluorophenyl)-5-isoxazolyl]ethenyl]-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 145911-07-5 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[2-[3-(3-methylphenyl)-5-isoxazolyl]ethenyl]-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 145911-08-6 ZCAPLUS

CN Benzeneacetic acid, 2-[2-[3-(4-chlorophenyl)-5-isoxazolyl]ethenyl]- α -ethylidene-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

RN 145911-09-7 ZCAPLUS

CN Benzeneacetic acid, 2-[2-[4-chloro-3-(3-chlorophenyl)-5-isoxazolyl]ethenyl]- α -ethylidene-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 145911-10-0 ZCAPLUS

CN Benzeneacetic acid, 2-[2-[1-(4-chlorophenyl)-1H-pyrazol-4-yl]ethenyl]- $\alpha-\text{ethylidene-}$, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 145911-49-5 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[(1-phenyl-1H-pyrazol-4-yl)oxy]methyl]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

RN 145911-50-8 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[[1-(4-methylphenyl)-1H-pyrazol-4-yl]oxy]methyl]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 145911-51-9 ZCAPLUS

CN Benzeneacetic acid, α -ethylidene-2-[[[1-(4-fluorophenyl)-1H-pyrazol-4-yl]oxy]methyl]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 145911-70-2 ZCAPLUS

CN Benzeneacetic acid, $2-[[[3-(4-\text{chlorophenyl})-5-\text{isoxazolyl}] \text{oxy}] \text{methyl}] - \alpha-\text{ethylidene-, methyl ester, (E)- (9CI)}$ (CA INDEX NAME)

145911-74-6 ZCAPLUS

CN Benzeneacetic acid, 2-[[[1-(4-cyanophenyl)-1H-pyrazol-4-yl]oxy]methyl]- α -ethylidene-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

L89 ANSWER 79 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1993:73650 ZCAPLUS Full-text

DOCUMENT NUMBER: 118:73650

TITLE: Antimycotic phenylacetic acid derivatives INVENTOR(S): Sauter, Hubert; Lorenz, Gisela; Steiner, Gerd; Janssen, Bernd; Anke, Timm; Steglich, Wolfgang

BASF A.-G., Germany PATENT ASSIGNEE(S): SOURCE: Eur. Pat. Appl., 20 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE	
EP 515901	A1	19921202	EP 1992-108035		19920513 <	
R: AT,	BE, CH, DE, DI	K, ES, FR,	GB, IT, LI, NL, PT,	SE		
DE 4117371	A1	19921203	DE 1991-4117371		19910528 <	
CA 2069691	A1	19921129	CA 1992-2069691		19920527 <	
JP 05170648	A	19930709	JP 1992-134851		19920527 <	
US 5334607	А	19940802	US 1992-889418		19920528 <	
PRIORITY APPLN. I	NFO.:		DE 1991-4117371	A	19910528	
OTHER SOURCE(S):	MARPA:	Γ 118:7365	0			

GI

$$\mathsf{MeXCOC} \bigvee_{V} \mathsf{V}$$

The phenylacetic acid derivs. I (X = 0, NH; Y = CHOMe, CHMe, CHEt, CHSMe, NOMe; Z = halo, NO2, CN, (un)substituted alkyl, aralkyl, aryloxyalkyl, etc.; U, V, W = H, Z, etc.) are medical fungicides. (E)-I (Z = 2-MeC6H4OCH2, X = 0, Y = CHOMe, U = V = W = H) had a min. inhibitory concentration of 0.1 μ g/mL against Aspergillus niger.

IT 145849-22-5 145849-23-6

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study); USES (Uses)

(fungicide, medical)

RN 145849-22-5 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[2-(3-phenyl-5-isoxazolyl)ethenyl]-, methyl ester, (E,?)- (9CI) (CA INDEX NAME)

Double bond geometry as described by E or Z.

RN 145849-23-6 ZCAPLUS

CN Benzeneacetic acid, α -(methoxymethylene)-2-[2-(1-phenyl-1H-pyrazol-4-yl)ethenyl]-, methyl ester, (E,?)- (9CI) (CA INDEX NAME)

Double bond geometry as described by ${\tt E}$ or ${\tt Z}$.

ACCESSION NUMBER: 1993:70035 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 118:70035

TITLE: Silver halide photographic material

INVENTOR(S): Okawa, Atsuhiro; Hirano, Shigeo; Obayashi, Keiji;

Ichijima, Yasushi

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 29 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 04248547	A	19920904	JP 1991-33463	19910204 <
PRIORITY APPLN. INFO.:			JP 1991-33463	19910204

AB The title material contains a compound represented by X(T)mY(ZPUG)4n [X = oxidation-reduction group; upon oxidation or reduction of X, the bond between X and (T)m is cleaved; T = linking group; Y = N-containing heterocyclic ring are given; Z = methylene (which is linked to a carbon atom of the said heterocyclic ring); PUG = photog. useful group; m = 0 or 1; n = 1 to 3]. The title material gives sharp images.

IT 145601-01-0

RL: TEM (Technical or engineered material use); USES (Uses) (photog. material containing)

RN 145601-01-0 ZCAPLUS

CN 1H-Tetrazole-1-acetic acid, 5-[[[1-[4-(hexadecylthio)-2,5-dihydroxyphenyl]-3,5-dimethyl-1H-pyrazol-4-yl]methyl]thio]-, butyl ester (9CI) (CA INDEX NAME)

L89 ANSWER 81 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1992:184505 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 116:184505

TITLE: Silver halide photographic material

INVENTOR(S): Ohashi, Hirobumi; Kawashima, Yasuhiko; Kagawa, Nobuaki

PATENT ASSIGNEE(S): Konica Co., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 28 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
JP 03204640	A	19910906	JP 1990-386	19900108 <	
PRIORITY APPLN. INFO.:			JP 1990-386	19900108	
GI					

- AB The title material on a support has at least one layer containing a dispersion of solid particles of a pyrazolone oxonol dye I (R1 = a substituent; R2 = H, alkyl, alkenyl, cycloalkyl, etc.; L1-L3 = a methine linkage; E = an acidic ring needed for forming an oxonol dye; n = 0-2). The title material shows excellent storage stability.
- RN 140214-21-7 ZCAPLUS
 CN 3-Pyridineacetic acid, 5-[[1-(4-carboxyphenyl)-5-hydroxy-3-methoxy-1H-pyrazol-4-yl]methylene]-1-ethyl-1,2,5,6-tetrahydro-2,6-dioxo-(9CI) (CA INDEX NAME)

- RN 140214-35-3 ZCAPLUS
- CN 1H-Pyrazole-1-propanoic acid, 4-[[1-[4-(aminosulfonyl)phenyl]-2-methyl-3,5-dioxo-4-pyrazolidinylidene]methyl]-5-hydroxy-3-methyl- (9CI) (CA INDEX NAME)

$$H_{2N}$$
 H_{2N}
 H

RN 140214-41-1 ZCAPLUS

CN 3-Pyridineacetic acid, 5-[3-[3-carboxy-1-(4-carboxyphenyl)-5-hydroxy-1H-pyrazol-4-yl]-2-propenylidene]-1,2,5,6-tetrahydro-4-methyl-2,6-dioxo-(9CI) (CA INDEX NAME)

PAGE 2-A

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L89 ANSWER 82 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1992:72186 ZCAPLUS Full-text

DOCUMENT NUMBER: 116:72186

TITLE: Silver halide photographic material INVENTOR(S): Yoshida, Kazuhiro; Hirabayashi, Kazuhiko

PATENT ASSIGNEE(S): Konica Co., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 18 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Fatent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND DATE		APPLICATION NO.	DATE	
JP 03223843 PRIORITY APPLN. INFO.:	А	19911002	JP 1990-20164 JP 1990-20164	19900130 < 19900130	
GI					

$$\begin{array}{c} \mathbb{R}^1 \\ \mathbb{N} \\ \mathbb{N} \\ \mathbb{R}^3 \end{array} \text{CH} \\ \mathbb{N} $

AB At least one layer of the title material contain dyes I (R1, R2 = carboxy, alkyl, aryl, alkoxycarbonyl, aryloxycarbonyl; R3-4 = sulfo- or carboxy- substituted alkyl or aryl) and an anionic surfactant, and is hardened by a hardening agent CH2:CHSO2(CH2)mO(LO)p(CH2)nSO2CH:CH2 (L = divalent organic group; m, n > 0; p = 0, 1). This photog. material provides low stain and high scratch resistance under rapid processing, and have high resistance to blocking by adhesion and high storage stability.

IT 138371-40-1

RL: USES (Uses)

(dye, backcoating of photog. films containing)

RN 138371-40-1 ZCAPLUS

CN 1H-Pyrazole-1-acetic acid, 4,5-dihydro-4-[[3-ethyl-5-hydroxy-1-(4-sulfophenyl)-1H-pyrazol-4-yl]methylene]-3-methyl-5-oxo-, disodium salt (9CI) (CA INDEX NAME)

ACCESSION NUMBER: 1991:559133 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 115:159133

TITLE: Preparation of pyrazolyl-substituted methyl

methoxyacrylates as agrochemical fungicides

INVENTOR(S): Oda, Masatsugu; Sakaki, Toshiro; Kikutake, Kuzuhiko

PATENT ASSIGNEE(S): Mitsubishi Kasei Corp., Japan

SOURCE: Eur. Pat. Appl., 76 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
	EP 433899	A1	19910626	EP 1990-124128	-	19901213 <
	EP 433899	B1	19950412			
	R: AT, BE, CH,	DE, ES	, FR, GB, II	r, LI, LU, NL		
	JP 04217668	A	19920807	JP 1990-324113		19901127 <
	JP 3018490	B2	20000313			
	CA 2031974	A1	19910614	CA 1990-2031974		19901211 <
	US 5055477	A	19911008	US 1990-625762		19901213 <
	AT 121080	T	19950415	AT 1990-124128		19901213 <
	ES 2074113	Т3	19950901	ES 1990-124128		19901213 <
	KR 157319	В1	19981116	KR 1990-20508		19901213 <
	US 5128481	A	19920707	US 1991-734292		19910717 <
PRIOF	ITY APPLN. INFO.:			JP 1989-323035	Α	19891213
				JP 1990-79763	Α	19900328
				JP 1990-273724	A	19901012
				JP 1990-324113	Α	19901127
				US 1990-625762	А3	19901213
001100	COLLDON (C)		115 150100			

OTHER SOURCE(S): MARPAT 115:159133

GΙ

R2 OCH2A

N C (= CHOMe)R

I Me C (X) n

Q1= (X) n

Q2= (X) m

$$Q2=$$
 (X) m

AB Pyrazolyl-substituted Me methoxyacrylates and analogs I (R1,R2 = H, C1-5 alkyl; A = Q1,Q2; X = H, halo, cyano, nitro, C1-10 alkyl, C1-10 alkoxy, etc.; m = 1,2; n = 1-5; R = CO2Me, cyano) were prepared Thus Et 4-benzyloxy-1,3-dimethylpyrazol-5-carboxylate (prepared by O-benzylation of Et 1,3-dimethyl-4-hydroxypyrazol-5-carboxylate) was reduced to the corresponding alc. by LiAlH4. This was converted to the chloride by SOC12, which was treated with NaCN to give (4-benzyloxy-1,3-dimethylpyrazol-5-yl)acetonitrile. Addition of the latter to a cooled solution of concentrated H2SO4 in MeOH gave the corresponding Me acetate derivative, which was condensed with HCOCO2Me, then

treated with Me2SO4 to give title compound II. An aqueous solution of II (200 ppm, stem-foliar application) gave 95% control of Puccina recondita on wheat.

IT 136193-00-5P 136193-01-6P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as agricultural fungicide)

RN 136193-00-5 ZCAPLUS

CN 1H-Pyrazole-5-acetic acid, α -(methoxymethylene)-1,3-dimethyl-4-[(2-phenyl-4-thiazolyl)methoxy]-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 136193-01-6 ZCAPLUS

CN 1H-Pyrazole-5-acetic acid, α -(methoxymethylene)-1,3-dimethyl-4-[(2-phenyl-4-thiazolyl)methoxy]-, methyl ester, (Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

L89 ANSWER 84 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1991:449697 ZCAPLUS Full-text

DOCUMENT NUMBER: 115:49697

TITLE: Preparation of 1-carbamoyl-3-(arylmethylthio)-1,2,4-

triazoles and S-oxidized analogs as herbicides

INVENTOR(S): Jelich, Klaus; Schmidt, Robert R.; Santel, Hans

Joachim; Luerssen, Klaus

PATENT ASSIGNEE(S): Bayer A.-G., Germany SOURCE: Ger. Offen., 21 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Fatent
LANGUAGE: German
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
				_	
DE 3929673	A1	19910314	DE 1989-3929673		19890907 <
EP 422369	A2	19910417	EP 1990-116317		19900825 <
EP 422369	A3	19920226			
R: BE, CH, DE,	FR, GB	, IT, LI, NL			
JP 03099066	A	19910424	JP 1990-233423		19900905 <
PRIORITY APPLN. INFO.:			DE 1989-3929673	Α	19890907
OTHER SOURCE(S):	CASREA	CT 115:49697	; MARPAT 115:49697		
GT					

- Title compds. [I; R1,R2 = C1-6 alkyl; R3 = 4-ClC6H4, cyanophenyl, nitrophenyl, (substituted) 5- or 6-membered heteroaryl, benzoxazolyl, benzothiazolyl; n = 0-2], were prepared as herbicides (no data). Thus, Et2NCOCl was added to a mixture of 3-(4-chlorobenzylthio)-2H-1,2,4-triazole (preparation given) in pyridine and the mixture was stirred 15 h to give 85% carbamoylated product, which was S-oxidized with 3-ClC6H4C(0)OOH in CHCl3 to give 79.5% title compound II. II was said to be very well tolerated by rice while showing good herbicidal activity.
- IT 134795-55-4P 134795-64-5P

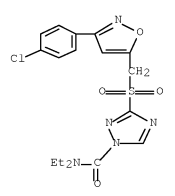
 RL: AGR (Agricultural use); BAC (Biological a

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as herbicide)

- RN 134795-55-4 ZCAPLUS
- CN 1H-1,2,4-Triazole-1-carboxamide, 3-[[[3-(4-chlorophenyl)-5-isoxazolyl]methyl]thio]-N,N-diethyl- (9CI) (CA INDEX NAME)

RN 134795-64-5 ZCAPLUS

CN 1H-1,2,4-Triazole-1-carboxamide, 3-[[[3-(4-chlorophenyl)-5-isoxazolyl]methyl]sulfonyl]-N,N-diethyl- (9CI) (CA INDEX NAME)



L89 ANSWER 85 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1990:552433 ZCAPLUS Full-text

DOCUMENT NUMBER: 113:152433

TITLE: Preparation of heterocyclic-substituted

 α -arylacrylates as pesticides and fungicides

INVENTOR(S): Schuetz, Franz; Neubauer, Hans Juergen; Kuekenhoehner,

Thomas; Schirmer, Ulrich; Hofmeister, Peter; Kuenast,

Christoph; Ammermann, Eberhard; Lorenz, Gisela;

Kardorff, Uwe

PATENT ASSIGNEE(S): BASF A.-G., Germany

SOURCE: Ger. Offen., 25 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Fatent
LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA]	CENT	NO.			KINI	D DATE	API	PLICATION NO.	DATE	
DE	3836	581			A1	19900503	DE	1988-3836581	19881027	<
CA	2000	362			A1	19900427	CA	1989-2000362	19891010	<
CA	2000	362			С	20010821				
CS	2744	76			В2	19910411	CS	1989-5825	19891013	<
ΙL	9198	8			Α	19930708	IL	1989-91988	19891013	<
ΕP	3787	55			A1	19900725	EP	1989-119384	19891019	<
ΕP	3787	55			В1	19931229				
	R:	AT,	BE,	CH,	DE,	ES, FR, GB,	GR, I	I, LI, NL, SE		
ΑT	9929	4			Τ	19940115	AT	1989-119384	19891019	<
ES	2061	878			Т3	19941216	ES	1989-119384	19891019	<
DD	2847	98			A5	19901128	DD	1989-333900	19891025	<
AU	8943	732			А	19900503	AU	1989-43732	19891026	<
AU	6211	56			В2	19920305				
ZA	8908	114			А	19910626	ZA	1989-8114	19891026	<
HU	2032	69			В	19910729	HU	1989-5455	19891026	<
JΡ	0218	0866			А	19900713	JP	1989-278765	19891027	<
KR	1277	69			В1	19980401	KR	1989-15489	19891027	<

US 5166216	A	19921124	US	1991-701019		19910513	<
US 5250553	A	19931005	US	1992-921765		19920730	<
US 5294628	A	19940315	US	1993-94580		19930716	<
US 5366984	A	19941122	US	1993-160836		19931203	<
PRIORITY APPLN. INFO.:			DE	1988-3836581	Α	19881027	
			US	1989-418664	В1	19891010	
			EP	1989-119384	Α	19891019	
			US	1991-701019	А3	19910513	
			US	1992-921765	А3	19920730	
			US	1993-94580	А3	19930716	
OTHER SOURCE(S):	CASREA	ACT 113:15243	33; 1	MARPAT 113:152433			

GΙ

AΒ Title compds. I (R = alkyl, alkenyl, haloalkyl, cycloalkyl, alkoxy, alkylcarbonyl, alkoxycarbonyl, halo, (substituted) aryl; Met = (N-Mesubstituted) 5-membered heteroarom. group containing 1-3 of O, S, and/or N and bound to A at a C atom; A = CH:CH, CH2CH2, CH2O, CH2S] were prepared as insecticides, acaricides, and nematocides (no data), and especially as fungicides for plants and materials. For example, Wittig-type reaction of di-Et 3-cyclopropylisoxazol-5-ylmethanephosphonate with Me 2-formylphenylacetate (preparation given) gave 47% Me 2-[2-(3- cyclopropylisoxazole-5yl)ethenyl]phenylacetate, which underwent condensation with Me formate (82%) and subsequent O-methylation of the resultant β -hydroxyacrylate (80%) to give title compound II. As a 0.05 weight% spray on grapevine leaves, II gave 100% protection against Plasmopara viticola.

ΙT 129562-62-5P 129590-29-0P

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction of, in preparation of heterocyclic-substituted arylacrylate fungicides)

129562-62-5 ZCAPLUS RN

CN Benzeneacetic acid, 2-[2-(3-cyclopropyl-5-isoxazolyl)ethenyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 129590-29-0 ZCAPLUS

CN Benzeneacetic acid, $2-[2-(3-\text{cyclopropyl}-5-\text{isoxazolyl})\text{ ethenyl}]-\alpha-(\text{hydroxymethylene})-, methyl ester (9CI) (CA INDEX NAME)$

IT 129562-60-3P 129562-61-4P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as fungicide and pesticide)

RN 129562-60-3 ZCAPLUS

CN Benzeneacetic acid, $2-[2-(3-\text{cyclopropyl}-5-\text{isoxazolyl})\text{ ethenyl}]-\alpha-$ (methoxymethylene)-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 129562-61-4 ZCAPLUS

CN Benzeneacetic acid, $2-[2-[3-(4-\text{chlorophenyl})-5-\text{isoxazolyl}]\text{ethenyl}]-\alpha-$ (methoxymethylene)-, methyl ester, (E,E)- (9CI) (CA INDEX NAME)

L89 ANSWER 86 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1990:188900 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 112:188900

TITLE: Silver halide photographic material containing oxonol

dye

INVENTOR(S): Kagawa, Nobuaki; Kawashima, Yasuhiko; Tanaka, Mari

PATENT ASSIGNEE(S): Konica Co., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 20 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Fatent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND DATE		APPLICATION NO.	DATE		
JP 01224749	А	19890907	JP 1988-50789	19880304 <		
PRIORITY APPLN. INFO.:			JP 1988-50789	19880304		
GI						

- AB In the title photog. material, ≥ 1 of photog. constitutional layers contains an oxonol dye (I) [R = cyano, R1CO, S02R1 (R1 = alkyl, aryl, heterocyclyl); J = divalent organic group; Z = CONR2, NR2CO, S02NR2, NR2SO2, CO2, OCO, S02, S02O, OSO2, NR2CONR3, O(CpH2qO)n, NR2CO2, OCONR2, NR2, SO, (R2, R3 = H, alkyl, aryl, heterocyclyl; p, q = 2-4; n \geq 1); sol = water-soluble functional group, or organic moiety with ≥ 1 of water-soluble functional groups; E = acid nucleus necessary to form an oxonol dye; L1-L5 = methine group; i, j, m = 0-1]. The dye is useful as filter dye, or in halation prevention or irradiation prevention.
- IT 126484-69-3
 - RL: USES (Uses)
 - (photog. antihalation dye)
- RN 126484-69-3 ZCAPLUS
- CN 1(2H)-Pyridineacetic acid, 5-cyano-3-[3-[3-cyano-1-(2,5-disulfophenyl)-5-hydroxy-1H-pyrazol-4-yl]-2-propenylidene]-3,6-dihydro-4-methyl-2,6-dioxo-, dipotassium salt (9CI) (CA INDEX NAME)

L89 ANSWER 87 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1990:148984 ZCAPLUS Full-text

DOCUMENT NUMBER: 112:148984

TITLE: Silver halide photographic material containing an

oxonol dye to prevent loss of image sharpness due to

halation

INVENTOR(S): Kagawa, Nobuaki; Kawashima, Yasuhiko; Tanaka, Mari

PATENT ASSIGNEE(S): Konica Co., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 17 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 01147452	A	19890609	JP 1987-307209	19871203 <
PRIORITY APPLN. INFO.:			JP 1987-307209	19871203
GI				

$$\mathbb{RC}(:0) \xrightarrow{\mathbb{N}} \mathbb{L}(\mathbb{L}^{1} = \mathbb{L}^{2})_{1}(\mathbb{L}^{3} = \mathbb{L}^{4})_{m} \xrightarrow{\mathbb{N}} \mathbb{R}^{3}$$

AB The photog. material having ≥1 hydrophilic colloid layer(s) contains in ≥1 of its component layer(s) an oxonol dye I (R = alkyl, aryl, heterocyclic group; R1 = alkyl, aryl, heterocyclic group substituted by sulfo, carboxyl or their salt; R2 = H, alkyl, aryl, heterocyclic ring; L, L1-4 = methyne; R3 = alkyl, aryl, heterocyclic ring, carboxyl, alkoxyl, aryloxy, carbamoyl, amino, acylamino, imido, ureido, hydroxy, cyano, alkoxycarbonyl, aryloxycarbonyl; l, m = 0, 1). It has an effective spectral filtering or antihalation function,

does not affect the photog. properties and is easily washed out during processing. Thus, in the manufacturing of a multilayer color paper, dye I (R, R3 = Me; R1, R2 = 2,5-di-sulfophenyl(K salt); l = 1; m = 0), dye I (R = Me; R3 = C02Et; R1 = 2,5-di-sulfophenyl(K salt); R2 = 4-sulfophenyl(K salt); l = 1; m = 0) and dye I (R = Me; R3 = CN; R1 = 2,5-di-sulfophenyl(K salt); R2 = 4-sulfophenyl(K salt); l = 1; m = 1) were added to green-sensitive layer, interlayer and red-sensitive layer resp.

IT 125367-70-6

RL: USES (Uses)

(antihalation dye, for photog. paper)

RN 125367-70-6 ZCAPLUS

CN 1H-Pyrazole-3-carboxylic acid, 4-[[1-(2,5-disulfophenyl)-1,5-dihydro-5-oxo-3-(1-oxopropyl)-4H-pyrazol-4-ylidene]methyl]-5-hydroxy-1-(2-sulfoethyl)-, tripotassium salt (9CI) (CA INDEX NAME)

●3 K

L89 ANSWER 88 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1989:574091 ZCAPLUS Full-text

DOCUMENT NUMBER: 111:174091

TITLE: 4-[(Isoxazolyl or styryl)methylene]thiohydantoin

derivatives as aldose reductase inhibitors

INVENTOR(S): Ogawa, Kazuo; Yamawaki, Ichiro; Matsushita, Yoichi

PATENT ASSIGNEE(S): Taiho Pharmaceutical Co., Ltd., Japan

SOURCE: PCT Int. Appl., 36 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:	TENT NO	Э.			KINI)	DATE		AP	PLICA:	TION NO	•		DATE	
WO	890289				A1	_	1989	0406	WO	1988-	-JP979			19880927	<
	W: 2 RW: 2	,	•		DE,	FR,	GB,	IT,	LU, N	L, SE					
JP	011569	965			A		1989	0620	JP	1988-	-187252			19880726	<
AU	88248	42			А		1989	0418	AU	1988-	-24842			19880927	<
PRIORIT	Y APPLI	N	INFO	.:					JP	1987-	-245591		A	19870929	
									JP	1988-	-187252		A	19880726	
									WO	1988-	-JP979	1	A	19880927	

OTHER SOURCE(S): MARPAT 111:174091

$$R^1CH$$
 $NR4$
 $Q=R2$
 N
 N

AB The title compds. [I; R1 = (tetrahydro)benzoisoxazolyl, (α-methyl)styryl, Q; R2 = halo, lower alkyl, CF3, MeO, phenethyl, PhCH2O, EtO2C, cyclopropyl, isobutylcyclohexyl, cyclohexylmethoxy, (halo or methoxy)phenyl, tetrahydropyranyl, thienyl, pyridyl); R3, R4 = H, lower alkyl, HO2CCH2, halobenzyl] were prepared as aldose reductase inhibitors. A mixture of 5-propylisoxazol-3-aldehyde, 2-thiohydantoin-1-acetic acid, NaOAc, Ac2O, and AcOH was refluxed 3 h to give 59% I (R1 = 5-n-propylisoxazol-3-yl, R3 = H, R4 = CH2CO2H). I inhibited aldose reductase prepared from a supernatant liquid of homogenized rats' crystalline lenses and 1M phosphate buffer (pH 6.2) with IC50's of 2.5-12 + 10-8M. Tablets (300 mg) were formulated from I (R1 = 5-tert-butylisoxazol-3-yl, R3 = H, R4 = CH2CO2H) 100, lactose 47, corn starch 50, crystalline cellulose 50, hydroxypropylcellulose 15, talc 2, magnesium stearate 2, ethylcellulose 30, unsatd. aliphatic acid glyceride 2 and TiO2 2 mg.

IT 122817-01-0P 122817-03-2P 122817-08-7P 122817-10-1P 122817-13-4P 122817-24-7P 122817-30-5P 122829-12-3P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as aldose reductase inhibitor)

RN 122817-01-0 ZCAPLUS

CN 1-Imidazolidineacetic acid, 4-[[3-(4-methoxyphenyl)-5-isoxazolyl]methylene]-5-oxo-2-thioxo-(9CI) (CA INDEX NAME)

RN 122817-03-2 ZCAPLUS

CN 1-Imidazolidineacetic acid, 5-oxo-4-[[3-(3-pyridinyl)-5-isoxazolyl]methylene]-2-thioxo-(9CI) (CA INDEX NAME)

RN 122817-08-7 ZCAPLUS

CN 1-Imidazolidineacetic acid, 4-[[3-(4-chlorophenyl)-5-isoxazolyl]methylene]-5-oxo-2-thioxo-(9CI) (CA INDEX NAME)

RN 122817-10-1 ZCAPLUS

CN 1-Imidazolidineacetic acid, 5-[[3-(4-chlorophenyl)-5-isoxazolyl]methylene]-4-oxo-2-thioxo-(9CI) (CA INDEX NAME)

RN 122817-13-4 ZCAPLUS

CN 1-Imidazolidineacetic acid, 4-[[5-[4-(2-methylpropyl)cyclohexyl]-3-isoxazolyl]methylene]-5-oxo-2-thioxo- (9CI) (CA INDEX NAME)

RN 122817-24-7 ZCAPLUS

CN 1-Imidazolidineacetic acid, 5-oxo-4-[[5-(2-thienyl)-3-isoxazolyl]methylene]-2-thioxo- (9CI) (CA INDEX NAME)

RN 122817-30-5 ZCAPLUS

CN 1-Imidazolidineacetic acid, 5-oxo-4-[[3-(tetrahydro-2H-pyran-2-yl)-5-isoxazolyl]methylene]-2-thioxo- (9CI) (CA INDEX NAME)

RN 122829-12-3 ZCAPLUS

CN 1-Imidazolidineacetic acid, 4-[(5-cyclopropyl-3-isoxazolyl)methylene]-5-oxo-2-thioxo-(9CI) (CA INDEX NAME)

$$0 \xrightarrow{N} CH \xrightarrow{H} S$$

$$CH_2 - CO_2H$$

L89 ANSWER 89 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1987:5026 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 106:5026

TITLE: [(Pyrazolylalkoxy)phenyl]ureas

INVENTOR(S): Go, Atsushi; Usui, Yoshihiro; Endo, Keiji; Hikido,

Mitsuru

PATENT ASSIGNEE(S): Mitsubishi Petrochemical Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 11 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Fatent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 61197559 PRIORITY APPLN. INFO.: GI	А	19860901	JP 1985-37127 JP 1985-37127	19850226 < 19850226

- The title compds. [I; R = alkyl, halo, CF3, Ph; n = 0-3; R1 = Me, MeO, H; R2 = H, halo, CF3; X = CH2, CH2CH2, CHMe], useful as herbicides, were prepared Thus, 3,4-Cl(HO)C6H3NHCONMe2 in CH2Cl2 containing NaH was treated with 1- (chloromethyl)pyrazole-HCl at room temperature for 2.5 h to give II. II was almost 100% effective against Echinochloa crus-galli at 10 kg/ha.
- IT 105675-70-5P

 RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as herbicide)
- RN 105675-70-5 ZCAPLUS
- CN Urea, N'-[3-chloro-4-[(5-methyl-3-phenyl-1H-pyrazol-1-yl)methoxy]phenyl]-N,N-dimethyl- (9CI) (CA INDEX NAME)

Ph
$$\sim$$
 NH \sim N

L89 ANSWER 90 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1986:424270 ZCAPLUS Full-text

DOCUMENT NUMBER: 105:24270

TITLE: Herbicidal thiadiazolylureas

INVENTOR(S): Morland, Robert B.; Cooke, Anson R.; Bishop, John R.

PATENT ASSIGNEE(S): Union Carbide Corp., USA

SOURCE: U.S., 18 pp.

CODEN: USXXAM

DOCUMENT TYPE: Fatent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

GΙ

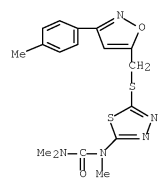
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 4576629	A	19860318	US 1984-589724	19840315 <
PRIORITY APPLN. INFO.:			US 1984-589724	19840315
OTHER SOURCE(S):	CASREA	ACT 105:24270); MARPAT 105:24270	

$$R(CR^1R^2)_{nS}(0)_{m}$$
 $N-N$ $N-N$ $NR^3CONR^4R^5$

- The title compds. I (R = heterocyclic or fused heterocyclic group; R1 and R2 are H, alkyl, cycloalkyl, alkoxy, carbalkoxy, halo; n = 1, 2, 3, 4, 5; m = 0, 1, 2; R3, R4, and R5 are H, cycloalkyl, Ph, alkyl, alkoxy), which showed herbicidal activity, were prepared A 2-amino-5-mercapto-1,3,4- thiadiazole was etherified and then treated with 1,1'-carbonyldiimidazole and Me2NH to give I (R = 2-thienyl, R1 = R2 = H, n = 1, m = 0, R3 = R4 = R5 = Me).

 II 102902-11-49 102902-12-59
- RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as herbicide)
- RN 102902-11-4 ZCAPLUS
- CN Urea, [5-[[[3-(4-chlorophenyl)-5-isoxazolyl]methyl]thio]-1,3,4-thiadiazol-2-yl]trimethyl- (9CI) (CA INDEX NAME)

- RN 102902-12-5 ZCAPLUS
- CN Urea, trimethyl[5-[[[3-(4-methylphenyl)-5-isoxazolyl]methyl]thio]-1,3,4-thiadiazol-2-yl]- (9CI) (CA INDEX NAME)



L89 ANSWER 91 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1986:50811 ZCAPLUS Full-text

DOCUMENT NUMBER: 104:50811

TITLE: Metalation of isoxazolyloxazolines, a facile route to

functionally complex isoxazoles: utility, scope, and

comparison to dianion methodology

AUTHOR(S): Natale, Nicholas R.; McKenna, John I.; Niou, Chorng

Shyr; Borth, Mark; Hope, Hakon

CORPORATE SOURCE: Dep. Chem., Univ. Idaho, Moscow, ID, 83843, USA

SOURCE: Journal of Organic Chemistry (1985), 50(26),

5660-6

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 104:50811

GΙ

$$\bigvee_{Me}^{R}\bigvee_{Me}^{Me}$$

AB 2-(5'-Alkyl-4'-isoxazolyl)- Δ 2-oxazolines I were lithiated at the C-5' alkyl group, and the lithio anions quenched with alkyl halides, aldehydes, and acylpyridinium salts as electrophiles. The lithio anion was also oxygenated with, e.g., N-(phenylsulfonyl)oxaziridine. The isoxazolyloxazoline system was converted into an isoxazolecarboxylic acid, an aldehyde, a ketone, and a chiral oxazoline. I were formed, metalated, and deprotected in synthetically useful yields, and represented a facile entry into functionally complex isoxazoles. To determine the necessity of the oxazoline protection/deprotection scheme, dianions of isoxazole-4- carboxylic acids were studied. The dianion method was found to be more efficient for simple alkyl halides, but limited in scope.

IT 99298-97-2P 99298-98-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

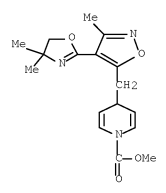
(preparation and oxidation of)

RN 99298-97-2 ZCAPLUS

CN 1(2H)-Pyridinecarboxylic acid, 2-[[4-(4,5-dihydro-4,4-dimethyl-2-oxazolyl)-3-methyl-5-isoxazolyl]methyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 99298-98-3 ZCAPLUS

CN 1(4H)-Pyridinecarboxylic acid, 4-[[4-(4,5-dihydro-4,4-dimethyl-2-oxazolyl)-3-methyl-5-isoxazolyl]methyl]-, methyl ester (9CI) (CA INDEX NAME)



L89 ANSWER 92 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1985:24226 ZCAPLUS Full-text

DOCUMENT NUMBER: 102:24226

TITLE: Contributions to the chemistry of tetraketones, III.

Synthesis and some reactions of 1,6-bis(p-

hydroxyphenyl)-1,3,4,6-hexanetetrone

AUTHOR(S): Kovac, Spomenka; Rapic, Vladimir; Lacan, Marijan CORPORATE SOURCE: Fak. Nahrungsmitteltechnol., Univ. Osijek, Osijek,

YU-54000, Yugoslavia

SOURCE: Liebigs Annalen der Chemie (1984), (10),

1755-8

CODEN: LACHDL; ISSN: 0170-2041

DOCUMENT TYPE: Journal LANGUAGE: German

OTHER SOURCE(S): CASREACT 102:24226

GΙ

AB Condensation of 4-HOC6H4COMe with (CO2Et)2 gave the title compound, which, e.g., with o-C6H4(NH2)2 gave the quinoxaline I and with PhNHNH2 gave the bipyrazole II.

IT 93846-85-6P

RN 93846-85-6 ZCAPLUS

CN 1,3-Propanedione, 1-[4-[[(methylamino)carbonyl]oxy]phenyl]-3-[5-[4-[[(methylamino)carbonyl]oxy]phenyl]-1H-pyrazol-3-yl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{HN} \\ \text{N} \\ \text{CH}_2 \\ \text{O} \\ \text{C} \\ \text{NHMe} \\ \text{MeNH} \\ \text{C} \\ \text{O} \\ \text{C} \\ \text{NHMe} \\ \text{MeNH} \\ \text{C} \\ \text{O} \\ \text{C} \\ \text{NHMe} \\ \text{MeNH} \\ \text{C} \\ \text{O} \\ \text{C} \\ \text{NHMe} \\ \text{NH} \\ \text{C} \\ \text{NHMe} \\ \text{NH} \\ \text{C} \\ \text{NHMe} \\ \text{NH} \\ \text{NH} \\ \text{C} \\ \text{NH} \\ \text{NH} \\ \text{C} \\ \text{NH} \\ \text{NH} \\ \text{NH} \\ \text{C} \\ \text{NH} \\ \text{NH} \\ \text{C} \\ \text{NH} \\ \text{NH} \\ \text{NH} \\ \text{C} \\ \text{NH} \\ \text{NH} \\ \text{C} \\ \text{NH} \\ \text{NH} \\ \text{NH} \\ \text{NH} \\ \text{C} \\ \text{NH} \\ \text{NH} \\ \text{NH} \\ \text{C} \\ \text{NH} \\ \text$$

L89 ANSWER 93 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1984:591912 ZCAPLUS Full-text

DOCUMENT NUMBER: 101:191912

TITLE: Substituted 4-imidazolyl pyrazoles with

antithromboembolic action

INVENTOR(S): Elbe, Hans Ludwig; Perzborn, Elisabeth; Seuter,

Friedel

PATENT ASSIGNEE(S): Bayer A.-G., Fed. Rep. Ger.

SOURCE: Ger. Offen., 45 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Fatent
LANGUAGE: German
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 3300795	A1	19840712	DE 1983-3300795	19830112 <
EP 115640	A2	19840815	EP 1983-113222	19831230 <
R: AT, BE, CH,	DE, FR	, GB, IT, LI	, LU, NL, SE	

JP 59130881
PRIORITY APPLN. INFO.:
OTHER SOURCE(S):

A 19840727 JP 1984-2126 DE 1983-3300795

19840111 <--A 19830112

CASREACT 101:191912; MARPAT 101:191912

GΙ

$$\mathbb{R}^{1}$$
 \mathbb{N}
 - AB The title compds. [I; R = H, alkyl, alkenyl, alkynyl, alkanoyl, alkoxycarbonyl, (un)substituted Ph, PhCH2, heteroaryl; R1 = (un)substituted alkyl, cycloalkyl] were prepared Thus, 1-(1H-imidazol-1-yl)-3, 3-dimethyl-2-butanone was condensed with Me2NCH(OMe)2 to give 90.5% pentenone II. II was cyclocondensed with N2H4 to give 55.3% I (R = H, R1 = Me3C) (III). III inhibited the aggregation of blood platelets with a min. inhibitory concentration of 1+10-5-3+10-5 g/mL.
- IT 92782-09-7P 92782-16-6P

RN 92782-09-7 ZCAPLUS

CN Benzoic acid, 4-[2-[4-(1H-imidazol-1-yl)-1H-pyrazol-3-yl]-2-methylpropoxy]- (9CI) (CA INDEX NAME)

RN 92782-16-6 ZCAPLUS

CN Benzoic acid, 4-[2-[4-(1H-imidazol-1-yl)-1H-pyrazol-3-yl]-2-methylpropoxy]-, methyl ester (9CI) (CA INDEX NAME)

L89 ANSWER 94 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1983:505236 ZCAPLUS Full-text DOCUMENT NUMBER: 99:105236

TITLE: Urea derivatives and their use for controlling

undesired plant growth

INVENTOR(S): Becker, Rainer; Theobald, Hans; Schirmer, Ulrich;

Spiegler, Wolfgang; Seufert, Walter; Wuerzer, Bruno

PATENT ASSIGNEE(S): BASF A.-G., Fed. Rep. Ger.

SOURCE: Ger. Offen., 52 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

GΙ

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
DE 3148291	A1	19830609	DE 1981-3148291		19811205 <
IL 67286	A	19860331	IL 1982-67286		19821117 <
CA 1187887	A1	19850528	CA 1982-415913		19821118 <
US 4500340	A	19850219	US 1982-443523		19821122 <
EP 81141	A1	19830615	EP 1982-110858		19821124 <
EP 81141	В1	19850731			
R: AT, BE, CH	, DE, FI	R, GB, IT,	LI, NL		
AT 14577	Τ	19850815	AT 1982-110858		19821124 <
JP 58113177	A	19830705	JP 1982-207783		19821129 <
BR 8207050	A	19831011	BR 1982-7050		19821203 <
ZA 8208894	A	19831026	ZA 1982-8894		19821203 <
PRIORITY APPLN. INFO.:			DE 1981-3148291	Α	19811205
			EP 1982-110858	А	19821124
OTHER SOURCE(S):	CASRE	ACT 99:1052	236; MARPAT 99:105236		

RZOn NHCONR²Me I Me CH₂O
$$\mathbb{R}^3$$
 II

AB Herbicidal (no data) I [R = (un)substituted isoxazolyl, benzothiazolyl, oxadiazolyl, etc.; R1 = H, Me, F3C, halo; R2 = H, alkyl, alkenyl, alkynyl, alkoxy; Z = alkylene; n = 0, 1] were prepared Thus, 113 g 3-methyl-5-isoxazolemethanol was treated with 4-FC6H4NO2 to give 201 g II (R3 = NO2), which (220 g) was reduced with SnCl2 to give 139 g II (R3 = NH2). This (20.4 g) was acylated with ClCONMe2 to give 14.1 g II (R3 = NHCONMe2).

IT 86913-13-5P 86913-15-7P 86913-23-7P
RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 86913-13-5 ZCAPLUS

CN Urea, N'-[4-[[3-(4-chlorophenyl)-5-isoxazolyl]methoxy]phenyl]-N,N-dimethyl-(9CI) (CA INDEX NAME)

RN 86913-15-7 ZCAPLUS
CN Urea, N,N-dimethyl-N'-[4-[(3-phenyl-5-isoxazolyl)methoxy]phenyl]- (9CI)
(CA INDEX NAME)

RN 86913-23-7 ZCAPLUS

CN Urea, N'-[4-[[3-(4-chlorophenyl)-5-isoxazolyl]methoxy]phenyl]-N-methoxy-N-methyl- (9CI) (CA INDEX NAME)

DOCUMENT NUMBER: 94:47172

TITLE: Simple synthesis of 4-(heteroarylmethyl)phenols and

their acylation

AUTHOR(S): Kuebel, Boerries

CORPORATE SOURCE: Hoechst A.-G., Frankfurt/Main, D-6230/80, Fed. Rep.

Ger.

SOURCE: Liebigs Annalen der Chemie (1980), (9),

1392-401

CODEN: LACHDL; ISSN: 0170-2041

DOCUMENT TYPE: Journal LANGUAGE: German

OTHER SOURCE(S): CASREACT 94:47172

GI

HO

$$CH_2$$
 R_1
 R_1
 R_1
 R_1
 R_2
 R_1
 R_2
 R_1
 R_2
 R_1
 R_2
 R_1
 R_2
 R_1

AB Condensation of 4-HOC6H4CHO with MeCOCH2COR (R = OEt, Me) gave 4-HOC6H4CH:C(COR)COMe, which were hydrogenated to 4-HOC6H4CH2CH(COR)COMe (I). These reacted with hydrazines or HONH2 to give II [R1 = Me, OH; X = NR2 (R2 = H, Me, m-tolyl, 3-ClC6H4), O]. Reaction of I (R = OEt) with acetamidine gave III. On acylation with MeNCO, II (R1 = Me, X = NH) and III are esterified at the phenolic OH group, whereas acid chlorides reacted with the R1 group of II (R1 = OH). In the case of II (R1 = Me, X = NH), the tendency toward O- or N-acylation depended on the base used.

IT 75999-10-9P 75999-20-1P

RL: SPN (Synthetic preparation); PREP (Preparation)

III

(preparation of)

RN 75999-10-9 ZCAPLUS

CN Phenol, 4-[[1-(3-chlorophenyl)-3,5-dimethyl-1H-pyrazol-4-yl]methyl]-, methylcarbamate (ester) (9CI) (CA INDEX NAME)

RN 75999-20-1 ZCAPLUS

CN 1H-Pyrazol-5-ol, 1-(3-chlorophenyl)-3-methyl-4-[[4-[[(methylamino)carbonyl]oxy]phenyl]methyl]- (9CI) (CA INDEX NAME)

L89 ANSWER 96 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1977:453160 ZCAPLUS Full-text

DOCUMENT NUMBER: 87:53160

TITLE: Heterobicyclics; Part IV. Imidazole N-oxides. VII.

Imidazo[4,5-c]pyrazoles from 4-nitro-5-

benzylaminopyrazoles

AUTHOR(S): Lange, Marina; Quell, Ruediger; Lettau, Herbert;

Schubert, Hermann

CORPORATE SOURCE: Sekt. Chem., Martin-Luther-Univ. Halle-Wittenberg,

Halle/Saale, Ger. Dem. Rep.

SOURCE: Zeitschrift fuer Chemie (1977), 17(3), 94-5

CODEN: ZECEAL; ISSN: 0044-2402

DOCUMENT TYPE: Journal LANGUAGE: German

OTHER SOURCE(S): CASREACT 87:53160

GΙ



AB Imidazopyrazoles I [R = Ph, 4-MeC6H4, 4-ClC6H4, 4-BrC6H4, 4-H2NC6H4, 3-H02CC6H4, 2,4-Me2C6H3, 2,4-H0(O2N)C6H3] were prepared by cyclizing II with base and reducing the 4-oxides of I with P(OEt)3 or TiCl3. II were obtained by aminating the 5-chloropyrazole. II (R = 2-hydroxy-1-naphthyl) did not cyclize.

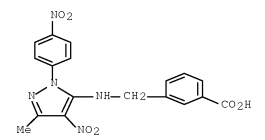
IT 63451-62-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and cyclization of)

RN 63451-62-7 ZCAPLUS

CN Benzoic acid, 3-[[[3-methyl-4-nitro-1-(4-nitrophenyl)-1H-pyrazol-5-yl]amino]methyl]- (9CI) (CA INDEX NAME)



L89 ANSWER 97 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1976:421354 ZCAPLUS Full-text

DOCUMENT NUMBER: 85:21354

TITLE: Substituted 4-[pyrazolyl-(1)-methylene]

oxazolin-5-ones

INVENTOR(S): Vogel, Christian; Braeuniger, Harald; Kristen, Helmut;

Peseke, Klaus

PATENT ASSIGNEE(S): Ger. Dem. Rep. SOURCE: Ger. (East), 3 pp.

CODEN: GEXXA8

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DD 117228	A1	19760105	DD 1975-183770	19750123 <
PRIORITY APPLN. INFO	0.:		DD 1975-183770	A1 19750123

Eto2C NNCH OPh

AB The pyrazolylmethyleneoxazolinone I was obtained in 45% yield by treating 4-ethoxymethyleneoxazolinone with H2NNHCH:C(CN)CO2Et.

IT 59681-38-8P

RN 59681-38-8 ZCAPLUS

CN 1H-Pyrazole-4-carboxylic acid, 5-amino-1-[(5-oxo-2-phenyl-4(5H)-oxazolylidene)methyl]-, ethyl ester (9CI) (CA INDEX NAME)

L89 ANSWER 98 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1974:82795 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 80:82795

TITLE: Syntheses of pyrazolone and pyrazole derivatives.

III. Syntheses of 3-substituted 5-methoxy-1-

phenylpyrazole derivatives

AUTHOR(S): Izumi, Kihathiro; Kitamikado, Tadashi; Sugiura, Shoji;

Kato, Kazuo; Hori, Mikio; Fujimura, Hajime

CORPORATE SOURCE: Res. Lab., Maruko Seiyaku Co., Ltd., Kasugai, Japan

SOURCE: Yakugaku Zasshi (1973), 93(10), 1349-55

CODEN: YKKZAJ; ISSN: 0031-6903

DOCUMENT TYPE: Journal LANGUAGE: Japanese

GI For diagram(s), see printed CA Issue.

AB For studies on biological activity, syntheses of 3-substituted 5-methoxy-1-phenylpyrazoles were attempted. N-Substituted 5-methoxy-1-phenylpyrazol-3-ylacetamides I (R = H2NCO, MeNHCO, EtNHCO, etc.) were obtained from I (R = CO2H, CO2Et, COCl). 3-Amino-methyl-5- methoxy-1-phenylpyrazole (II) was synthesized from I (R = CO2H or H2NNHCO) by the Schmidt reaction or Curtius reaction. N-Acylmethylamine derivs., e.g. I (R = o-H2NC6H4CONH) and Melubrin-type I (R = NaO3SCH2NH) were obtained from II, but the attempt to prepare N-alkylamine derivs. of II was unsuccessful. 3-Chloromethyl-5-methoxy-1-phenylpyrazole was synthesized from II, and N-alkylamine derivs., e.g. I (R = MeNH), and sulpyrin type I (R = NaO3SCH2NMe) were obtained from I (R = C1).

IT 51862-40-9P

RN 51862-40-9 ZCAPLUS

CN Benzoic acid, 2-[[(5-methoxy-1-phenyl-1H-pyrazol-3-yl)acetyl]amino]-, methyl ester (9CI) (CA INDEX NAME)

L89 ANSWER 99 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1967:465090 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 67:65090

ORIGINAL REFERENCE NO.: 67:12303a,12306a

TITLE: Pyrazolone stabilizers for poly- α -olefins

INVENTOR(S): Harris, Raymond Clement; Newland, Gordon C.

PATENT ASSIGNEE(S): Eastman Kodak Co.

SOURCE: U.S., 6 pp. CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

GI For diagram(s), see printed CA Issue.

The stabilizers have the general formula I. Thus, low-d. polyethylene (II) of melt index 2 was compounded (roll temps. 220°F. and 270°F.) with 1% I (R1 = R4 = Ph, R2 = R3 = Me) for 4 min. and molded into 125-mil thick sheets. When exposed, under stress, to natural weathering and to a Twin-Arc Weather-Ometer, >24 months and 3000 hrs., resp., were required before cracking occurred, compared with 12 months and 330 hrs. for control (II with no additive). No color change occurred. I (R1 = Ph, R2 = Me, R3 = iso-Bu, R4 = H) was used similarly to stabilize II. Similarly used to stabilize polypropylene (III), a 15:85 butene-propylene copolymer, and a 20:80 ethylene-propylene copolymer were the following I (R1-4, resp., given): Ph, Me, Me, Ph; Ph, Me, iso-Bu, H; Ph, Me, NH2, Ph; Ph, Me, CF3, H; Ph, Me, CO2H, Ph. Also used to stabilize III were the following I (R1-4, resp., given): Ph, Ph, iso-Bu, H; Ph, Me, Ph, H; Ph, Me, Ph, CH2CH2OH; Ph, Me, Me, SO2Ph; Ph, Me, Me, p-O2NC6H4; Ph, Me, CO2Et, Ph; Ph, Me, CO2Et, H.

IT 18468-43-4

RL: USES (Uses)

(as ultraviolet stabilizer for olefin polymers)

RN 18468-43-4 ZCAPLUS

CN Pyrazole-3-carboxylic acid, 5-hydroxy-4-[(3-methyl-5-oxo-1-phenyl-2-pyrazolin-4-ylidene)methyl]-, ethyl ester (8CI) (CA INDEX NAME)

L89 ANSWER 100 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1967:104945 ZCAPLUS Full-text

DOCUMENT NUMBER: 66:104945

ORIGINAL REFERENCE NO.: 66:19627a,19630a

TITLE: Some glyoxals with isoxazole and oxazole nuclei and

their derivatives

AUTHOR(S): Giannella, M.; Gualtieri, Fulvio CORPORATE SOURCE: Univ. Camerino, Camerino, Italy

SOURCE: Bollettino Chimico Farmaceutico (1966), 105,

708-18

CODEN: BCFAAI; ISSN: 0006-6648

DOCUMENT TYPE: Journal LANGUAGE: Italian

GI For diagram(s), see printed CA Issue.

AΒ The title compds. are prepared and tested as antivirals and bacteriostatics. Thus, 23 g. 2,5-dimethyl-4-acetyloxazole in dioxane is added to 21.97 g. SeO2 in 275 ml. dioxane and 27.5 ml. H2O, the solution refluxed 20 hrs., boiled with C, and filtered, and the solvent evaporated in vacuo to give 28 g. I (R = COCH(OH)2), m. $112-14^{\circ}$ (C6H6). The following I are prepared (R and m.p. given): COCH:NOH, 170-70.5°; COCH:NNHCONH2, 219° (decomposition); COCH: NNHCSNH2, 217-18° (decomposition); COCH: NC6H4CO2H-p, 159-60°; COCH:NC6H4SO2NH2-p, 219-20°; COCH:NNHCOC5H4N-4, 206-8°; C(CH:NNHCONH2):NNHCONH2, 238° (decomposition); C(CH:NNHCOC5H4N-4):NNHCOC5H4N-4, 241-2°; 5-Phenyl-3-acetylisoxazole (2 g.) is added to 1.42 g. SeO2 in 30 ml. 10% H2O-dioxane. After the theoretical amount of Se is collected, the same volume of H2O is added, the mixture boiled with C and filtered, and the solvent evaporated in vacuo to give II (R = COCH(OH)2), m. $55-6^{\circ}$ (H2O). The following II are prepared (R and m.p. given): COCH:NNHCONH2, 200-1°; COCH:NNHCSNH2, 224-5° (decomposition); COCH:NC6H4CO2H-p, 99-100°; COCH:NNHCOC5H4N-4, 203-4°. 3-Phenyl-4-acetyl-5-methylisoxazole (2 g.) in dioxane is added to 1.32 g. SeO2 in 20 ml. 10% H2O-dioxane, the mixture refluxed until the separation of Se is complete, boiled with C and filtered and the solvent evaporated in vacuo to give 2.5 g. III (R = COCH(OH)2) m. 87-9° (H2O). The following III are prepared (R and m.p. given): COCH:NOH, 143-4°; COCH:NNHCONH2, 196-7° (decomposition); COCH:NNHCSNH2, 173-4°; COCH:NC6H4CO2H-p (IV), 132-3°; COCH:NNHCOC5H4N-4 (V), 165-6°; Glyoxal monohydrate (0.5 q.) in EtOH is treated with 0.35 q. concentrated aqueous KOH, the solution heated 8 hrs. and cooled, H2O added, and the mixture acidified with dilute HCl (3-phenyl-5-methyl-4-isoxazolylhydroxyacetic acid, m. 87.8°(water). Formaldehyde (20 ml., 40%) and 40 ml. 15% NH3 are added slowly with stirring and cooling to 1.0 g. glyoxal monohydrate in 20 ml. EtOH and the mixture kept overnight and treated with ice to give 4-(3-phenyl-5-methyl- 4isoxazolyl)imidazole, m. 219-20° (HCONMe2). Acetylacetone (6 q.) is added slowly, dropwise with stirring into a EtONa solution (obtained from 1.37 g. Na and 100 ml. absolute EtOH) and the mixture cooled with water-ice mixture, treated with 12 g. p-nitrobenzohydroxamoyl chloride in 100 ml. absolute EtOH, kept overnight, and filtered to give 10.4 g. 3-(p-nitrophenyl)-4- acetyl-5methylisoxazole (VI), m. $147-8^{\circ}$ (EtOH). VI (7.5 g.) in dioxane is added to 4.05 g. SeO2 in 54 ml. dioxane-6 ml. H2O. Refluxing 24 hrs. Se separation, water addition, boiling with C, filtration, and solvent elimination gave 9.5 g. VII (R = COCH(OH)2), m. $108-9^{\circ}$ (H2O). The following are prepared (R and m.p. given): COCH:NOH, 180-1°; COCH:NNHCONH2, 219-20° (decomposition); COCH: NNHCSNH2, 201-2° (decomposition); CHCH: NC6H4CO2H-p, 214-15°; COCH:NNHCOC5H4N-4, 223-4.5°; COCH:NC(:NH)NH2, 185-7°. V (2 q.) in 80 ml. hot EtOH is boiled 1 hr. with the same volume of 10% K2CO3 and the mixture cooled and acidified with dilute HCl to precipitate 5-(3-p-nitrophenyl-5-methyl-4isoxazoly1)-1,2,4-triazine-3-thione, m. 216° (HCONMe2-H2O). IV (2.5 g.) in 60ml. hot EtOH is boiled 1 hr. with 60 ml. 10% K2CO3, the solvent evaporated in vacuo, and the residue acidified to precipitate 5-(3-p-nitrophenyl-5-methyl-4isoxazolyl-)1,2,4-triazin-3-one, m. 241-2° (AcOH). Glyoxal monohydrate (1 g.) in EtOH is treated with an aqueous solution of 0.45 g. KOH and the mixture kept overnight at room temperature, treated with ice, and acidified with dilute HCl to precipitate (3-p-nitrophenyl-5- methyl-4isoxazolyl) hydroxyacetic acid, m. 159-60° (H2O). Glyoxal hydrate (1 g.) in 20 ml. EtOH is treated with 30 ml. 40% formaldehyde and slowly with stirring 50 ml. concentrated NH3 and the mixture kept overnight at room temperature, and diluted with water to precipitate 4-(3-p-nitrophenyl-5-methyl-4isoxazolylimidazole, m. 213-14° (HCONMe2-water). The most biol. active compound is IV.

13788-07-3F 13788-11-9F 13788-18-6F
RL: SPN (Synthetic preparation); PREP (Preparation)

ΙT

(preparation of) RN 13788-07-3 ZCAPLUS

CN Benzoic acid, p-[[[(5-phenyl-3-isoxazolyl)carbonyl]methylene]amino]- (8CI) (CA INDEX NAME)

RN 13788-11-9 ZCAPLUS

CN Benzoic acid, p-[[[(5-methyl-3-phenyl-4-isoxazolyl)carbonyl]methylene]amin o]- (8CI) (CA INDEX NAME)

RN 13788-18-6 ZCAPLUS

CN Benzoic acid, p-[[[[5-methyl-3-(p-nitrophenyl)-4-isoxazolyl]carbonyl]methylene]amino]- (8CI) (CA INDEX NAME)

L89 ANSWER 101 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1963:482229 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 59:82229
ORIGINAL REFERENCE NO.: 59:15267f-q

TITLE: Ability of the two methyl groups of the quaternary

base of 3,5-dimethylisoxazole to couple. IV. Syntheses

of diacylmethane derivatives.

AUTHOR(S): Lampe, W.; Smolinska, J.

CORPORATE SOURCE: Univ. Warsaw

SOURCE: Bulletin de l'Academie Polonaise des Sciences, Serie

des Sciences Chimiques (3963), 11(2), 49-53

CODEN: BAPCAQ; ISSN: 0001-4095

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

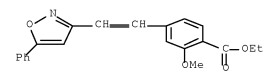
AB CA 53,5243d. 3-[(4-Hydroxy-3-methoxybenzylidene)methyl]-5-phenylisoxazole EtI salt was converted to its 4'-carbethoxy analog (I), m. 135°. I was reduced with H-Pt to 1-phenyl-5-(4-hydroxy-3-methoxyphenyl)-3- iminopentan-1-one (II), m. 136° (100% yield), which with concentrated HCl gave the corresponding 1,3-pentanedione (III) m. 73.5° (60% yield). The corresponding 3'-ethoxy analogs were prepared, from I (m. 152°) (50% yield) from II (m. 133.5°) (100% yield), and from III (m. 112°) (60% yield). Infrared data indicate no absorption at 1700-1 cm., hence β -imino- and β -diketones are enolized or have intermol. H-bonds

IT 94870-25-4P, o-Anisic acid, 4-[2-(5-phenyl-3-isoxazolyl)vinyl]-,

ethyl ester

RN 94870-25-4 ZCAPLUS

CN o-Anisic acid, 4-[2-(5-phenyl-3-isoxazolyl)vinyl]-, ethyl ester (7CI) (CA INDEX NAME)



L89 ANSWER 102 OF 102 ZCAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1910:17866 ZCAPLUS Full-text

DOCUMENT NUMBER: 4:17866

ORIGINAL REFERENCE NO.: 4:3196b-i,3197a-i,3198a

TITLE: Indigoid Dyes. VI. Aliphatic Aromatic Compounds

AUTHOR(S): Felix, A.; Fried-Lander, P.

SOURCE: Monatshefte fuer Chemie (1910), 31, 55-79

CODEN: MOCMB7; ISSN: 0026-9247

DOCUMENT TYPE: Journal
LANGUAGE: Unavailable
OTHER SOURCE(S): CASREACT 4:17866

AB The synthesis of various indigoid dyes is here described; the blue nuance of indigo is least changed when the NH group is substituted by CH : CH. When substituted by the CONH there is a shifting to blue-violet, by S to a carmine-red, and CH2 and CO to a red-orange; maxima of absorption (Angstrom units) have been determined for the following: Bis-2-indoleindigo $\lambda 6120$, 2-naphthalene-2-indoleindigo $\lambda 6440$ and 5910, 2-isoquinolone-2-indoleindigo

 λ 5840, 2-thionaphthene-2- indoleindigo λ 5790, 2-indane-2-indoleindigo λ 4920, 2-indanone-2-indoleindigo λ 3080. 2-Indane-2-indoleindigo (I), from equivalent parts ketohydrindene and isatin chloride, red needles; warming with H2SO4 gives a sulphonic acid. It is not attacked by NaOH (40%). Sublimes (decompose) in fine needles. 2-Indanone-2-indoleindigo (II), from equivalent parts diketohydrindene and α -isatin anilide, brownish violet needles. Boiling with dilute NaOH and addition of NaCl there separates, as a decompose product, the Na salt of 1-keto-3-hydroxyhydrindene-2-aldehyde, C10H6O3 (X). In the mother liquor anthranilic acid remains. The free aldehyde occurs in red needles, m. 139.5°. For the preparation of the coumaranone indigos dimethoxycoumaranone, C10H10O4, was first made by action of Me2SO4 and NaOH upon trihydroxychloroacetophenone, yellow needles m. 122.5-123°. When a dilute alkali and less Me2SO4 is used there results monomethyldihydroxycoumaranone, C9H8O4, needles, m. 197°. The former compound gives with Br a mono- and a dibromo derivative Warming equivalent parts of lphaisatin chloride and dimethoxycoumaranone yields a dimethoxycoumarane-2indoleindigo (III), copper-red crystals. The hydroxymethoxycoumaranone with isatin chloride gives a similar dye, C17H11O5N, more easily attacked by alkali than dye (III). According to J. Prochazka the dimethoxy and hydroxymethoxycoumaranone unite readily with aldehydes (in presence of acid or alkali) to form O isologs of indogenides. The combination with BzH yields the compound (XIV), pale yellow prisms, m. 148-9°. Combination with salicylaldehyde yields the compound (XV), light orange-yellow needles, m. 240°. Combination with m-hydroxybenzaldehyde yields the compound, C17H14O5 yellow needles m. 202.5-203°. Combination with p-hydroxybenzaldehyde yields the compound, C17H14O5, citron-yellow. The most colored here is the oderivative, the least the m-. Combination with protocatechuic aldehyde yields the compound (XVI), orange-yellow needles m. 217°. The dimethyl ether of this, m. 194-194.5°, results when piperonaldehyde is used. Hydroxymethoxycoumaranone condensed with piperonaldehyde yields an analogous compound C17H12O6, m. 190-190.5°. As with aldehyde so also dimethoxycoumaranone unites readily with β -naphthoguinone-4-sulphonic acid to form 2-hydroxynaphthalene-2-dimethoxycoumaranindolignone (XI), orange-brown needles, 1-oxy-3-isoquinoline-2-indoleindigo (IV), prepared from isatin chloride and dioxyisoquinoline. Dark blue needles. 1,3-Phenylmethyl-4pyrazole-2-indoleindigo (VII), from phenylmethylpyrazolone and isatin- α anilide, lustrous black plates, soluble without change in concentrate H2SO4 with red-brown color turning to a blue-red by dilution. The substance dissolves unacted upon in 10% NaOH solution but on boiling decomposes into anthranilic acid and the Na salt of 1,3-phenylmethyl-5- pyrazolone-4-aldehyde (IX), white needles, m. $173-4^{\circ}$. Of this aldehyde there was prepared the phenylhydrazone C17H16ON4, light yellow needles m. 159°; the aldazine C32H22O2N6 orange needles m. 290°. The aldehyde unites quantitatively with anthranilic acid giving an azomethin (XII), light yellow needles m. 240°. When equivalent parts 3-methyl-pyrazolone and isatin anilide are warmed in PhNO2 solution there results 3-methyl-4-pyrazole-2-indoleindigo (VI), dark violet needles, dissolving to a carmin in most solvents and readily attacked by alkali. The condensation of HSCN with lpha-isatin anilide in Ac2O gives 5thiazothiole-2-indoleindigo (V), black needles subliming to dark violet, 5thiazolthiole-2-thionaphtheneindigo (xIII) results from thioisatin anilide and HSCN, red-brown needles. Dioxypyrimidine-2- indoleindigo (VIII) results from condensation of barbituric acid with α -isatin anilide; small crystals of metallic luster.

861527-19-7P, Anthranilic acid, N-(5-hydroxy-3-methyl-1-phenyl-4-ΙT pyrazolylmethylene) -RL: PREP (Preparation)

(preparation of)

- RN 861527-19-7 ZCAPLUS
- CN Anthranilic acid, N-(5-hydroxy-3-methyl-1-phenyl-4-pyrazolylmethylene)-

=> d his full (FILE 'HOME' ENTERED AT 07:40:12 ON 28 SEP 2007) FILE 'ZCAPLUS' ENTERED AT 07:40:26 ON 28 SEP 2007 FILE 'REGISTRY' ENTERED AT 07:47:44 ON 28 SEP 2007 STRUCTURE UPLOADED L1L2 STRUCTURE UPLOADED L3 19 SEA SSS SAM L2 D SCA STRUCTURE UPLOADED T.4 50 SEA SSS SAM L2 AND L4 L5 FILE 'STNGUIDE' ENTERED AT 08:08:42 ON 28 SEP 2007 D STAT QUE L5 FILE 'REGISTRY' ENTERED AT 08:16:31 ON 28 SEP 2007 L*** DEL 22011 S L2 AND L4 FULL SSS D COST FULL D STAT QUE L6 L*** DEL 8082 S N2C3/ESS AND L6 STRUCTURE UPLOADED L6 50 SEA SSS SAM L2 AND L4 AND L6 L7 L8 SCREEN 1840 50 SEA SSS SAM L2 AND L4 AND L6 AND L8 L9 49 S L7 NOT L9 L*** DEL L*** DEL 49 S L9 NOT L7 1 S L9 NOT L11 L*** DEL D SCA 50 S L9 AND NRS>2 L*** DEL 50 S L7 AND NRS>2 L*** DEL D STAT QUE L9 STRUCTURE UPLOADED L10 L11 41 SEA SSS SAM L2 AND L10 AND L8 775523 SEA ABB=ON PLU=ON N2C3/ES OR NOC3/ES L12 30896 SEA ABB=ON PLU=ON NSC3/ES L13 805906 SEA ABB=ON PLU=ON L12 OR L13 464 SEA ABB=ON PLU=ON NPC3/ES L14L15 L16 806370 SEA ABB=ON PLU=ON (L13 OR L14 OR L15) 50 SEA SUB=L16 SSS SAM L2 L17 D L4 D STAT QUE L17 61080 SEA SUB=L16 SSS FUL L2 L18 SAVE TEMP L18 JAI214STR2BL/A L19 STRUCTURE UPLOADED L20 50 SEA SUB=L16 SSS SAM L19 L21 71084 SEA SUB=L16 SSS FUL L19 SAVE TEMP JAI214STR19B/A L21 FILE 'STNGUIDE' ENTERED AT 08:47:53 ON 28 SEP 2007 FILE 'REGISTRY' ENTERED AT 08:51:10 ON 28 SEP 2007 L22 STRUCTURE UPLOADED L23 STRUCTURE UPLOADED 50 SEA SUB=L21 SSS SAM L23 L24

D STAT QUE L24

31522 SEA SUB=L21 SSS FUL L23

L25

SAVE TEMP JAI214STR23B/A L25 L26 STRUCTURE UPLOADED FILE 'STNGUIDE' ENTERED AT 09:08:21 ON 28 SEP 2007 FILE 'REGISTRY' ENTERED AT 09:35:02 ON 28 SEP 2007 L27 STRUCTURE UPLOADED L28 50 SEA SUB=L25 SSS SAM L27 L29 STRUCTURE UPLOADED L30 50 SEA SUB=L25 SSS SAM L29 STRUCTURE UPLOADED L31 20 SEA SUB=L25 SSS SAM L31 L32 FILE 'STNGUIDE' ENTERED AT 09:54:10 ON 28 SEP 2007 FILE 'REGISTRY' ENTERED AT 09:56:28 ON 28 SEP 2007 L33 STRUCTURE UPLOADED L34 15 SEA SUB=L25 SSS SAM L33 D SCA L35 262 SEA SUB=L25 SSS FUL L33 SAVE TEMP L35 JAI214STR33L/A FILE 'ZCAPLUS' ENTERED AT 10:00:32 ON 28 SEP 2007 L36 82 SEA ABB=ON PLU=ON L35 FILE 'REGISTRY' ENTERED AT 10:00:47 ON 28 SEP 2007 FILE 'ZCAPLUS' ENTERED AT 10:02:22 ON 28 SEP 2007 E US2005-517214/APPS L37 1 SEA ABB=ON PLU=ON US2005-517214/AP D SCA SEL RN FILE 'REGISTRY' ENTERED AT 10:03:00 ON 28 SEP 2007 876 SEA ABB=ON PLU=ON (100-39-0/BI OR 100-51-6/BI OR 103324-26-1/ L38 BI OR 103626-03-5/BI OR 105170-18-1/BI OR 107-08-4/BI OR 107-18-6/BI OR 109492-77-5/BI OR 111196-81-7/BI OR 111493-88-0/ BI OR 1140-69-8/BI OR 114474-04-3/BI OR 116-53-0/BI OR 119-36-8/BI OR 123-25-1/BI OR 123374-28-7/BI OR 128796-39-4/BI OR 139-85-5/BI OR 140-88-5/BI OR 141-75-3/BI OR 14191-95-8/BI OR 14199-15-6/BI OR 1423-26-3/BI OR 1423-27-4/BI OR 148-53-8/BI OR 148872-79-1/BI OR 149490-75-5/BI OR 152270-53-6/BI OR 152468-10-5/BI OR 152608-83-8/BI OR 1556-18-9/BI OR 15802-80-9/ BI OR 15964-81-5/BI OR 15971-92-3/BI OR 16063-70-0/BI OR 160721-25-5/BI OR 16110-09-1/BI OR 167762-83-6/BI OR 1700-30-7/ BI OR 171817-14-4/BI OR 1722-10-7/BI OR 174607-36-4/BI OR 176214-15-6/BI OR 178547-21-2/BI OR 18368-64-4/BI OR 19438-10-9 /BI OR 2011-06-5/BI OR 20349-89-7/BI OR 20921-09-9/BI OR 20921-14-6/BI OR 209404-16-0/BI OR 20967-96-8/BI OR 212688-07-8 /BI OR 2150-44-9/BI OR 220380-56-3/BI OR 23795-02-0/BI OR 24214-73-1/BI OR 258506-68-2/BI OR 26691-25-8/BI OR 26691-27-0/ BI OR 26691-29-2/BI OR 27772-62-9/BI OR 29682-12-0/BI OR 32884-23-4/BI OR 32884-25-6/BI OR 328919-24-0/BI OR 33252-28-7/ BI OR 33577-16-1/BI OR 342023-83-0/BI OR 342023-88-5/BI OR 342023-90-9/BI OR 342024-10-6/BI OR 342024-14-0/BI OR 342024-99 -1/BI OR 342026-17-9/BI OR 35857-89-7/BI OR 367259-04-9/BI OR 36873-42-4/BI OR 372-48-5/BI OR 38275-43-3/BI OR 394-50-3/BI

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              O SEA ABB=ON PLU=ON L35 AND L38
L39
           379 SEA ABB=ON PLU=ON L25 AND L38
L40
L41
            50 SEA SUB=L25 SSS SAM L19
L42
             15 SEA SUB=L35 SSS SAM L19
                D STAT QUE L30
         16848 SEA SUB=L25 SSS FUL L29
L43
                SAVE TEMP L43 JAI214STR29B/A
                STRUCTURE UPLOADED
L44
L45
                STRUCTURE UPLOADED
L46
                STRUCTURE UPLOADED
L47
             50 SEA SUB=L43 SSS SAM L46
L48
           8395 SEA SUB=L43 SSS FUL L46
                SAVE TEMP L48 JAI214STR46B/A
L49
           3169 SEA ABB=ON PLU=ON L48 AND NRS<4
                E "PROPANOIC ACID, 2-METHYL-2-((4-(2-((3-METHYL-1-[4-(TRIFLUORO
              1 SEA ABB=ON PLU=ON "PROPANOIC ACID, 2-METHYL-2-((4-(2-((3-METH
L50
                YL-1-(4-(TRIFLUOROMETHOXY)PHENYL)-1H-PYRAZOL-5-YL)AMINO)ETHYL)-
                2-THIAZOLYL) THIO) -, MONOHYDROCHLORIDE"/CN
                D SCA
                D RSD
           1312 SEA ABB=ON PLU=ON L49 AND 16.165.12/RID
L51
            29 SEA ABB=ON PLU=ON L49 AND 16.299.11/RID
L52
            309 SEA ABB=ON PLU=ON L40 AND L51
L53
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L54
           326 SEA ABB=ON PLU=ON L51
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L*** DEL
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L55
           1569 SEA ABB=ON PLU=ON L48 AND NOC3/ES
                E "BENZOIC ACID, 4-((((5-METHYL-3-PHENYL-4-ISOXAZOLYL)CARBONYL)
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L56
                OXAZOLYL) CARBONYL) AMINO) METHYL) -, METHYL ESTER"/CN
                D RSD
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L57
           2091 SEA ABB=ON PLU=ON L51 OR L57 323 SEA ABB=ON PLU=ON L58 AND L38
L58
L59
            56 SEA ABB=ON PLU=ON L40 NOT L59
L60
             63 SEA ABB=ON PLU=ON NSC3/ES AND L48
L61
               E "BUTANOIC ACID, 2-METHYL-2-(4-((3-(4-(TRIFLUOROMETHYL)PHENYL)
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L62
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                D RSD
L63
              5 SEA ABB=ON PLU=ON 16.171.9/RID AND L49
                D SCA
L64
           2096 SEA ABB=ON PLU=ON L58 OR L63
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L65
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     FILE 'REGISTRY' ENTERED AT 10:59:55 ON 28 SEP 2007
     FILE 'ZCAPLUS' ENTERED AT 11:00:50 ON 28 SEP 2007
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L66
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L67
           275 SEA ABB=ON PLU=ON L65 AND P/DT
L68
            26 SEA ABB=ON PLU=ON L66 AND PY<2003
L69
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L70
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L71
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L73
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L74
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L75
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L76
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263 SEA ABB=ON PLU=ON ODAKA H?/AU
7435 SEA ABB=ON PLU=ON KIMURA H?/AU
L77
L78
L79
L80
            14 SEA ABB=ON PLU=ON MIZUFUNE H?/AU
           169 SEA ABB=ON PLU=ON FUKATSU K?/AU
              2 SEA ABB=ON PLU=ON L72 AND (L76 OR L77 OR L78 OR L79 OR L80
L82
                OR L81)
             11 SEA ABB=ON PLU=ON L76 AND (L77 OR L78 OR L79 OR L80 OR L81)
1 SEA ABB=ON PLU=ON L77 AND (L78 OR L79 OR L80 OR L81)
L83
L84
L85
             15 SEA ABB=ON PLU=ON L78 AND (L79 OR L80 OR L81)
             1 SEA ABB=ON PLU=ON L79 AND (L80 OR L81)
L86
             1 SEA ABB=ON PLU=ON L80 AND L81
L87
             20 SEA ABB=ON PLU=ON (L82 OR L83 OR L84 OR L85 OR L86 OR L87)
L88
                D SCA L82
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FILE 'REGISTRY' ENTERED AT 11:12:15 ON 28 SEP 2007

FILE 'ZCAPLUS' ENTERED AT 11:12:20 ON 28 SEP 2007

D STAT QUE L88

D IBIB ABS HITIND L88 1-20

FILE 'REGISTRY' ENTERED AT 11:13:54 ON 28 SEP 2007

FILE 'ZCAPLUS' ENTERED AT 11:13:57 ON 28 SEP 2007

D STAT QUE L72

102 SEA ABB=ON PLU=ON L72 NOT L88

D IBIB ABS HITSTR L89 1-102

FILE HOME

L89

FILE ZCAPLUS

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This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE REGISTRY

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 26 SEP 2007 HIGHEST RN 948239-70-1 DICTIONARY FILE UPDATES: 26 SEP 2007 HIGHEST RN 948239-70-1

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TSCA INFORMATION NOW CURRENT THROUGH June 29, 2007

Please note that search-term pricing does apply when conducting ${\tt SmartSELECT}$ searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

FILE STNGUIDE

FILE CONTAINS CURRENT INFORMATION.

LAST RELOADED: Sep 24, 2007 (20070924/UP).

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